Investigating the Role of the Inhibitor of Apoptosis Proteins (IAPs) in Metastasis Formation

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Investigating the Role of the Inhibitor of Apoptosis Proteins (IAPs) in Metastasis Formation

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“Knowledge is in the end based on acknowledgement”-Ludwig Wittgenstein.

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DECLARATION OF AUTHORSHIP

I hereby certify that the thesis I am submitting is my own original work, except for experiments and analyses listed below:

- *In vivo* experiments performed using MDA-MB231 cells subcutaneously engrafted in NOD/SCID mice were performed by Dr. Daniele Lecis and Dr. Giacomo Manenti.

- Gene expression profiling and bioinformatics analyses were performed in collaboration with the Functional Genomics and Bioinformatic core facility of Fondazione IRCCS Istituto Nazionale dei Tumori.

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1 ABSTRACT

Inhibitor of apoptosis proteins (IAPs) constitute a conserved family of molecules, which regulate both apoptosis and receptor signalling. They are often deregulated in cancer cells and represent potential targets for therapy. In my work, I investigated the effect of IAP inhibition in vivo to identify novel down-stream genes expressed in an IAP-dependent manner that could contribute to cancer aggressiveness. To this end, immunocompromised mice engrafted subcutaneously with the triple negative breast cancer (TNBC) cell line MDA-MB231 were treated with SM83, a Smac mimetic developed in our laboratory that acts as a pan-IAP inhibitor, and tumour nodules were profiled for gene expression. The analysis revealed that the inhibition of IAPs significantly reduces the expression of SNAI2, a zinc finger transcriptional repressor often associated with cancer aggressiveness, resistance to therapy and metastatic potential, especially in breast cancer. By testing several TNBC cell lines, I found that SNAI2 levels is promoted specifically by cellular IAP1 (cIAP1), and not by other IAPs, and that SM83-dependent down-regulation of SNAI2 reduces cancer cell motility. Accordingly, cIAP1 depletion blocks epidermal growth factor receptor (EGFR)-dependent activation of the mitogen-activated protein kinase (MAPK) pathway causing the reduction of SNAI2 transcription levels. The inhibition of EGFR signalling stems from the block of receptor signalling and from the down-regulation of its levels, but paradoxically the silencing of cIAP1 promotes EGFR stability rather than its degradation. Nonetheless, EGFR levels decrease upon cIAP1 silencing due to reduced NF-kB-dependent gene expression supporting the notion that cIAP1 controls EGFR in an opposite fashion, promoting its gene expression while causing its degradation. In conclusion, my work indicates that IAP-targeted therapy could contribute to
EGFR inhibition and to the reduction of its down-stream mediators. This approach could be particularly effective in tumours characterized by high levels of EGFR and SNAI2, such as TNBCs.
2 INTRODUCTION

2.1 Apoptosis: a process of programmed cell death

Apoptosis is the most studied type of programmed cell death; it is regulated by molecular pathways, which are evolutionarily conserved among the species (Degterev and Yuan, 2008). Apoptosis is crucial during embryonic development of multicellular organisms and, therefore, its deregulation is responsible for several diseases (Favaloro et al., 2012). For instance, excessive apoptosis leads to neurodegeneration (Mattson, 2000), while insufficient activation of the apoptotic machinery results in autoimmune disorders (Nagata, 2010) and many cancer types. Accordingly, pathological inhibition of apoptosis has been shown to favour cancer development. Therefore, several approaches have been designed to induce apoptosis or prevent its inhibition in tumours in order to eradicate them.

Resistance to apoptosis is one of the hallmarks of cancer, which were proposed by Hannah and Weinberg (Hanahan and Weinberg, 2011; Figure 2.1). In fact, aggressive tumours develop various strategies to escape the apoptotic process, and there is therefore the need to target the apoptotic blocks in order to promote the activity of therapeutic treatments, which usually rely on apoptotic activation to be effective (Gimenez-Bonafe et al., 2009).
Figure 2.1 - Hallmarks of cancer (Hanahan and Weinberg, 2011). The figure shows the hallmarks of cancer that characterize neoplastic cells compared to normal cells.

2.1.1 Mechanisms of apoptosis

Two main pathways constitute the apoptotic machinery: the extrinsic and the intrinsic apoptotic programs, which both depend on the balance between pro- and anti-apoptotic effectors. The extrinsic pathway senses and propagates death-inducing signals and soluble factors originated from the microenvironment outside the cells, while the intrinsic circuit is triggered by a variety of intra-cellular signals (Fulda and Debatin, 2006; Galluzzi et al., 2012). Importantly, these two pathways are tightly linked and culminate in the activation of normally inactive proteases (caspases-8 and -9, respectively), which, in turn, trigger the effector caspases (caspases-3, -6 and -7). The final step is then the execution phase of apoptosis, thus leading to the cell disruption.

Although different mechanisms regulate programmed cell death, all the apoptosis pathways converge in the activation of caspases, which are therefore executioners of
apoptosis (Figure 2.2). Caspases are conserved cysteine proteases that, in the absence of apoptosis signals, are expressed as inactive zymogens (pro-caspases) and, upon strictly controlled activation, acquire the capability to cleave their substrates after an aspartate residue (Thornberry and Lazebnik, 1998). Caspases are classified in initiator and effector depending whether they are apical activators of the apoptotic pathway or if they amplify an up-stream stimulus received by another caspase. Of note, initiator and effector caspases are characterized by a different structure. Initiator caspases display a long pro-domain that interacts with the up-stream adapter molecules through two different motifs: the caspase recruitment domain (CARD; caspase-2, -9) and the death effector domain (DED; caspase-8 and -10). On the contrary, a short pro-domain defines effector caspases (caspase-3, -6 and -7) and allows the cleavage of diverse substrates, which have already been processed by up-stream caspases, leading to the apoptosis demolition phase. A vast number of substrates can be cleaved by caspases (Fischer et al., 2003), such as nuclear lamin, inhibitor of caspase activated DNase (ICAD) and especially cleaved poly(adenosine diphosphate-ribose)polymerase (PARP), which, upon cleavage, is universally exploited as a marker for apoptotic cells (Nosseri et al., 1994).
Figure 2.2 - Human caspase-mediated apoptosis (Lamkanfi, 2011). Based on their mechanism of action, caspases are distinguished in initiator caspases and effector caspases. Initiator caspase-8 and -10 comprise a DED motif in their long pro-domain, while the other initiator caspases show the CARD motif.

Importantly, the caspase-mediated proteolytic process is accompanied by discrete morphological changes resulting from various events, including DNA fragmentation, chromatin condensation and nuclear remodelling (Figure 2.3). Interestingly, apoptotic cells, on the basis of morphological alterations, can be distinguished from cells dying by diverse types of death, such as necrosis, necroptosis and autophagy (Degterev and Yuan, 2008; Galluzzi et al., 2015).

Figure 2.3 - Morphological changes in apoptotic cells. Adapted from (Lamkanfi, 2011). Apoptosis is accompanied by a variety of biochemical and morphological features, including nuclear condensation and oligo-nucleosomal DNA fragmentation, shrinkage of the cell, cytoplasmic packing in apoptotic bodies and rapid phagocytosis by neighbouring cells. Cleaved PARP is commonly used as marker of apoptosis. All these features uniquely identify the apoptosis process.

2.1.2 The intrinsic apoptotic pathway

The activation of the intrinsic apoptosis pathway is mediated by diverse non-receptor-induced stimuli that produce intracellular signals leading to mitochondrial-
dependent death. In response to the mitochondrial outer membrane permeabilization (MOMP), cytochrome c is released from the inter-mitochondrial membrane space to the cytosol, where forms the apoptosome by interacting with the initiator caspase-9 and apoptotic protease activating factor 1 (APAF-1; Reubold and Eschenburg, 2012). This event is tightly controlled by the B-cell lymphoma-2 (BCL-2) family members (Figure 2.4), and, in particular, it is inhibited by the anti-apoptotic (BCL-2, BCL-XL and myeloid cell leukaemia 1-MCL-1) proteins, whereas it is promoted by the pro-apoptotic (BCL-2 antagonist killer 1 –BAK, BCL-2-associated X protein –BAX and BH3-only protein -BID) ones. Following apoptosome formation, caspase-9 is cleaved and activates the effector caspases-3, -6 and -7. These caspases, in turn, cleave diverse cellular substrates hence causing the biochemical and morphological changes, which are classically associated with the apoptotic phenotype (Danial and Korsmeyer, 2004).

Figure 2.4 - The intrinsic apoptosis signalling pathway (Ashkenazi et al., 2017). Cellular stress, either DNA damage or endoplasmic reticulum (ER) stress, induces the intrinsic apoptosis pathway with consequent
inhibition of the anti-apoptotic proteins, BCL-2, BCL-XL, BCL-W and MCL-1, and subsequent activation of the pro-apoptotic ones, BAK and BAX. This event results in MOMP, which allows the release of cytochrome c from the mitochondria. Once in the cytosol, cytochrome c forms a complex with pro-caspase-9 and APAF-1, thus activating the caspase-9. The latter promotes the induction of pro-caspase-3 and -7, and consequently cell death.

2.1.3 The extrinsic apoptotic pathway

The extrinsic apoptotic program, also named death receptor-dependent pathway, is activated by the binding of death ligands belonging to the tumour necrosis factor (TNF) superfamily to its cognate receptors, the TNF-Receptors (TNF-Rs), which are localized in the cell surface (Walczak, 2013). The stimulation of the TNF-Rs leads to rapid activation of the initiator caspase-8 and -10 (Barnhart et al., 2003; Dickens et al., 2012; Ganten et al., 2004), which are critically involved in the activation of apoptosis signalling. In details, upon binding to its ligand, the adaptor proteins Fas-associated protein with death domain (FADD) and TNF receptor-associated protein with death domain (TRADD) are recruited to the receptor (Figure 2.5). The formed complex, named death-inducing signalling complex (DISC; Scaffidi et al., 1998), allows pro-caspases-8 and -10 dimerization and activation, which results in the processing of their down-stream effector caspases-3, -6 and -7 (McIlwain et al., 2013).
Figure 2.5 - Overview of the apoptosis pathways (Fulda and Vucic, 2012). Both external and internal stimuli are responsible for apoptosis pathway activation. The extrinsic pathway is induced by binding of death ligands to the cognate receptors. Upon ligand binding, cytoplasmic adapter proteins are recruited, thus forming the DISC. This complex is responsible for the auto-catalytic activation of initiator caspases triggering to the execution phase of apoptosis. The cross-talk between extrinsic and intrinsic pathways occurs at the mitochondria. When mitochondria undergo MOMP, the apoptosome is formed and activates the initiator caspase-9 thus propagating the apoptotic process.

Although caspase-3 is capable of cleaving a number of substrates necessary for cell viability, the activation of caspase-8 and -3 is not always sufficient to kill cells. Cells are therefore divided in two groups: “type I” and “type II” (Ozoren and El-Deiry, 2002). While in “type I” cells the activation of the extrinsic pathway is sufficient to induce irreversible damage that crucially compromises cell viability, “type II” cells require the involvement of mitochondria to efficiently activate apoptosis (Figure 2.6). In these cells, caspase-8 cleaves BID, thereby generating the activated truncated form of BID, t-BID (Kantari and Walczak,
which leads to MOMP via the direct activation of BAX and BAK. This step is therefore in common between the extrinsic and intrinsic pathways and allows their cross-talk.

Figure 2.6 - Regulation of cell death by mitochondria (Tait and Green, 2010). The cross-talk between intrinsic and extrinsic apoptosis pathways is mediated by mitochondria and occurs after caspase-8-mediated cleavage and subsequent activation of BID (t-BID, hereafter). T-BID causes MOMP, which favours the release of various proteins from the mitochondrial intermembrane space, thus promoting caspase activation and apoptosis.

2.2 IAPs: Inhibitors of apoptosis proteins

Extrinsic and intrinsic apoptotic signals are controlled by the inhibitor of apoptosis protein (IAP) family, which consists of eight members, called X-linked IAP (XIAP), cIAP1, cIAP2, neuronal-IAP (NAIP), Survivin, Livin (ML-IAP), Apollon (BRUCE) and ILP-2 (Figure 2.7). Although IAPs were initially described as being negative regulators of apoptosis, later works have clarified their role also in controlling several other cell features and mechanisms (Rothe

2011), which leads to MOMP via the direct activation of BAX and BAK. This step is therefore in common between the extrinsic and intrinsic pathways and allows their cross-talk.
et al., 1995; Damgaard and Gyrd-Hansen, 2011). The activity on caspases has been confirmed only for 2-3 members of the family and it is likely to be restricted to limited settings (Choi et al., 2009; Eckelman et al., 2006). IAPs are ubiquitously expressed and their expression is deregulated in diseases and cancer cells, where they contribute to chemoresistance and poor prognosis (Gyrd-Hansen and Meier, 2010). For this reason, IAPs are considered potential targets in clinical oncology and many pharmacological approaches have been designed to inhibit the members of this family.

2.2.1 IAP structure and conserved domains

IAPs are characterized by the presence of one or three conserved motifs of about 70 amino acids, named baculoviral IAP repeat (BIR) domains (Birnbaum et al., 1994), originally identified in the baculoviral genome. Through this domain, IAPs, also called BIR-containing proteins (BIRCs), interact with other proteins. XIAP, cIAP1, and cIAP2 are among the most characterized IAPs and contain three BIR domains (Figure 2.7) that mediate protein-protein interactions. Moreover, XIAP and cIAP1/2 bear a domain named really interesting new gene (RING), which confers the E3 ubiquitin ligase activity that allows the binding with the E2 ubiquitin-conjugating enzymes and catalyzes the transfer of ubiquitin to a target substrate (Vaux and Silke, 2005). Therefore, the RING domain provides IAPs with the capability to polyubiquitinate proteins, such as caspases and many other substrates, and also to self-ubiquitinate. Ubiquitination can induce either proteasomal degradation or regulation of the target proteins in several signalling cascades (Chen, 2012). Other two additional domains identify some IAPs: a caspase-associated recruitment domain (CARD) and an ubiquitin-associated (UBA) domain. CARD regulates the E3 activity preventing RING domain...
dimerization, which is essential for cIAP1 to exert its activity (Lopez et al., 2011). While cIAPs affect the apoptotic process only indirectly, XIAP is considered as a direct inhibitor of caspases. In particular, the BIR3 domain of XIAP interacts with caspase-9, preventing its activation, whereas activated caspase-3 and -7 are sequestered after the interaction with the BIR2 and the up-stream linker region (Chai et al., 2001; Datta et al., 2000; Riedl et al., 2001).

Figure 2.7 - Structural domains of IAP family members. Adapted from (de Almagro and Vucic, 2012). The IAP family is constituted by eight members characterized by one or three BIR domains, typically located in the protein amino(N)-terminus. Several mammalian IAPs share the presence of the additional structural domain RING, which functions as an E3 ubiquitin ligase, while only cIAP1 and cIAP2 show the presence of CARD motif. UBA motif is shared by XIAP, cIAP1/2 and ILP-2. Survivin has a single BIR domain combined with a COOH-terminal alpha-helix coiled-coil domain. While, NAIP structure shows three BIR motifs together with the nucleotide-binding oligomerization (NACHT) and leucine-rich repeats (LRR) domains.

Based on the capability to regulate caspases, BIR domains can be divided into two classes, I and II, where only type II can interact to caspases, while type I is responsible for the binding with other proteins, including TRAF1 and 2 (Gyrd-Hansen and Meier, 2010). Notably, type II BIRs are characterized by a conserved groove, which allows the interaction with a
region within caspases, called IAP binding motif (IBM; Wu et al., 2000). An example of IBM is constituted by the AVPI (Ala-Val-Pro-Ile) sequence which is located at the N terminus of the second mitochondria-derived activator of caspase/direct inhibitor of apoptosis-binding protein with low pi (Smac/DIABLO), a mammalian natural antagonist of IAPs (Figure 2.8). Interestingly, after apoptotic stimuli, Smac/DIABLO is released from the mitochondria into the cytosol, where it binds to the XIAP-BIR3 domain thus relieving the inhibition of caspase-9 and eventually allowing the activation of the executioner caspases-3 and -7.

![Figure 2.8 - Smac/DIABLO: the IAP natural antagonist](Fulda, 2015). The binding of Smac/DIABLO to XIAP neutralizes XIAP-mediated caspase inhibition, resulting in apoptosis cell death.

### 2.2.2 The role of IAPs in cancer

The crucial role of IAPs in oncogenesis derives from their capability to promote resistance against cell death and control many survival pathways. Not surprisingly, IAPs have been found aberrantly expressed in diverse types of tumour (Che et al., 2012; Hundsdoerfer et al., 2010). However, the prognostic significance of IAP over-expression remains not
completely clear. This is the case of XIAP, whose expression levels correlate with disease severity and prognosis in acute myelogenous leukemia (AML) and renal cell carcinoma, but not in non-small cell lung carcinoma or cervical carcinoma (LaCasse et al., 2008). Deregulated expression of IAPs can be a result of different events, such as gene amplifications, mutations or deletions and chromosomal translocations. For instance, amplification of 11q21-q23, in which both cIAP1 and cIAP2 are located, has been found in several tumours, including, glioblastoma, hepatocellular carcinoma, esophageal carcinoma, cervical cancer, liver cancer, non-small-cell lung cancer (NSCLC), and pancreatic cancer (LaCasse et al., 2008). Moreover, trans-activation and post-translational events are often responsible for the increase of IAP expression in tumours. For example, the t(11;18)(q21;q21) translocation event has been found in mucosa-associated lymphoid tissue (MALT) lymphoma and causes the fusion of the BIR domains of cIAP2 with the paracaspase MALT1 (Rosebeck et al., 2011). As a consequence of this phenomenon, the constitutive activation of the NF-κB signalling pathway (Darding et al., 2011) occurs thereby promoting oncogenesis and tumour progression.

2.2.3 Smac mimetic-mediated targeting of IAPs

Based on the capability to prevent apoptosis, IAPs represent possible targets for cancer therapy and this leads to the development of small compounds, which are specific for IAP domains. Hence, the elucidation of the biochemical structures involved in the IAP–Smac/DIABLO interaction allowed the development of Smac-like molecules, called Smac mimetics (SMs). SMs resemble the N-terminal region of Smac/DIABLO and, in particular, mimic the IBM motif (Eckelman et al., 2008), which is directed against the XIAP BIR2- and BIR3-caspase interaction pocket, and prevent XIAP-mediated caspase inhibition. While
initially designed to target XIAP, SMs have been shown to affect cIAP1/2 E3 activity, driving their self-ubiquitination and proteasomal degradation (Figure 2.9).

![Diagram of cIAP activation](image.png)

**Figure 2.9 - SM-mediated activation of cIAPs** (Feltham et al., 2011). The binding of SMs with cIAPs promotes the RING dimerization and activates the cIAP E3 ligase function, self-ubiquitination and proteasomal degradation.

Notably, about 10-15 % of cancer cell lines are efficiently killed *in vitro* by SMs as standalones (Petersen et al., 2007; Figure 2.10) with low nanomolar EC50s, depending on the molecules used. In 2007, it has been reported by several groups that the cytotoxic effect of the SMs in monotherapy does not depend on their activity against XIAP, but it stems from a rapid SM-dependent depletion of cIAP1 and cIAP2. This event results in the autocrine production of TNF and subsequent cancer cell death (Petersen et al., 2007; Varfolomeev et al., 2007; Vince et al., 2007).
SMs are usually considered not toxic in normal tissues and the anti-tumour activity of several SMs has been tested in phase I/II clinical trials (Fulda, 2015). They were developed either as monovalent compounds containing one tetrapeptide mimicking endogenous Smac/DIABLO, or as bivalent molecules having two mimicking sequences connected together by a chemical linker. Since the natural Smac/DIABLO protein acts as a dimer, bivalent SMs were realized to enhance the binding affinity compared to a monomer Smac AVPI-mimicking peptide. Of note, the binding of the Smac/DIABLO AVPI motif is stronger towards BIR3 compared to BIR2, but the presence of two AVPI-like motifs ensures the formation of a very stable interaction between SMs and IAPs by targeting simultaneously two BIR domains within the same IAP (Lecis et al., 2012). As a consequence of this stable interaction (Cossu et al., 2009b; Cossu et al., 2012), dimeric SMs have been shown to be much more effective both in vitro and in vivo (Fulda, 2014b) compared to monomeric ones. In addition, dimeric compounds are more efficient in causing the depletion of cIAP1 and cIAP2 (Varfolomeev et al., 2007). However, dimeric SMs are less “drug-like” compared to monomeric ones from a pharmacological viewpoint (Fulda, 2015). In fact, being smaller molecules, monovalent SMs can...
even be endowed with a good oral bioavailability, which provides an important advantage for their clinical application. Conversely, the bioavailability of bivalent SMs is limited by their molecular weight that requires the employment of other administration routes, such as intravenous injection.

Unfortunately, the vast majority of cancer cells are usually resistant to SM employed in monotherapy, but the reason for this resistance is not fully understood. It has been reported that cIAP2 degradation occurs in a cIAP1 dependent manner (Darding et al., 2011). Upon SM treatment, cIAP2 is quickly replaced by newly synthesized protein that cannot be degraded anymore in the absence of cIAP1. Importantly, cIAP2 displays some common features with cIAP1 and its presence is sufficient to confer resistance to SM treatment (Darding et al., 2011). As SMs are very rarely cytotoxic in monotherapy, the efficacy should be enhanced by the combination with either death agonists, such as TNFα, TNF-related apoptosis-inducing ligand (TRAIL; Lecis et al., 2010), and FasL, or chemotherapy (Greer et al., 2011), resulting in synergistic activity (Moller et al., 2014). SMs have been reported to synergise with chemotherapeutic drugs in solid tumours and hematologic malignancies. However, some side effects and unwanted toxicity caused especially by immune system activation and cytokine production have been revealed by in vivo experiments and the first clinical trials (Amaravadi et al., 2011; Infante et al., 2014; Sikic et al., 2011).

SM efficacy has been tested in phase I/II clinical trials for solid and hematologic malignancies (Fulda, 2015), as monotherapy or in rational combinations with gemcitabine, daunorubicin, cytarabine, or Paclitaxel. Importantly, phase I trials indicate SM administration as a safety and well tolerated treatment, even if inducing an increase of cytokines in patients
(Fulda, 2014b). For instance, the up-regulation of inflammatory cytokines, that are NF-kB target genes, has been reported by Infante and colleagues after the administration of LCL161 for the treatment of human advanced solid tumours (Infante et al., 2014). In the attempt to reduce possible risks linked to SM administration, the new compound Birinapant has been designed with reduced affinity for XIAP, but still conserving the same effect on cIAP1 and cIAP2 (Obexer and Ausserlechner, 2014). In fact, it has been reported that SM-triggered cytokine release stems from the inhibition of XIAP. Intriguingly, Birinapant treatment does not promote TNF production and results well tolerated (Allensworth et al., 2013; Condon et al., 2014). The efficacy of this compound has been also assessed together with other chemotherapeutics to analyse whether some clinical benefits may be obtained from the combinations (Krepler et al., 2013). Among SMs, the two monovalent compounds: LCL161 (Novartis) and AT-406/Debio1143 (Ascenta Therapeutics; Figure 2.11) are currently tested in clinical trials, for the treatment of refractory multiple myeloma and advanced solid malignacies or metastatic NSCLC, respectively.
2.3 The TNF/TNF-R superfamily

As already mentioned, SM-mediated autocrine production of TNF occurs in response of clAP depletion leading to cell death. TNF is a member of a large family of cytokines that exert pleiotropic functions among whom regulation of the immune system, control of cell proliferation, differentiation and apoptosis (Ashkenazi and Dixit, 1998). TNF is expressed as a transmembrane protein, membrane TNF (mTNF), which can be shed by the metalloprotease
TNF-converting enzyme (TACE) into a soluble factor (sTNF). While mTNF can activate both TNF-R1 and TNF-R2, the soluble form fully activates only TNF-R1, despite the high binding affinity with TNF-R2 (Richter et al., 2012).

The members of the TNF-R superfamily are classified into two groups based on the presence of the death domain (DD; Walczak, 2013). Receptors belonging to the first group display an intracellular DD, which confers the capability to induce cell death. Conversely, members belonging to the second class lack this domain and cannot directly activate cell death, but need to interact with additional adaptor molecules, such as TNF-receptor-associating factors (TRAFs) through the receptor TRAF-interacting motif (TIM; Dempsey et al., 2003). The recruitment of adaptor proteins is crucial for the activation of the intracellular pathways triggered by the TNF superfamily ligands. Of note, the DDs are present only in a few adaptor proteins (MacEwan, 2002). In fact, TRADD and FADD exploit their DD to interact with the receptors and activate cell death, whilst other interactors, such as TRAFs, bind to the receptors either directly via the receptor TIM domain or indirectly by recruiting other adaptor proteins as intermediates (Hehlgans and Pfeffer, 2005). For its increased capability to induce cell death, TNF-R1 signalling has been studied more deeply than TNF-R2 in the cancer field. Therefore, several functions of the latter receptor still remain unclear.

2.3.1 **TNF-R1 complex and its signal transduction**

While the carboxyl (C)-terminal end of TNF-R1 is occupied by DD motif for the apoptosis signal transduction, the N-terminal region contains a cysteine-rich domain that favours the preassembly of the receptor into a trimeric complex (Hehlgans and Pfeffer, 2005). TNF-R1 is constitutively expressed in all tissue and can activate a variety of cellular
processes, including cell proliferation, apoptosis or necroptosis, based on the cellular context (Wajant et al., 2003). Even though TNF-R1 usually activate the nuclear factor κB (NF-κB) and mitogen-activated protein kinase (MAPK) signalling pathways, TNF-R1 activation can result in different scenarios determined by the formation of two diverse complexes, named Complex I and II (Figure 2.7; Micheau and Tschopp, 2003). In the first scenario, TNF-R1 trimerizes within a few minutes upon TNF stimulation and recruits TRADD to the DD. Then, additional mediators, namely receptor-interacting protein 1 (RIP1; Wong et al., 2010), TRAF2 and/or TRAF5, are added to the complex and, in turn, cIAP1 and cIAP2 are assembled. The activation of NF-κB signalling pathway occurs after the recruitment of NF-κB essential modulator (NEMO; also known as IKKγ), TGFβ-activated kinase 1 (TAK1; also known as MAP3K7) and the TAK1-binding proteins (TAB1 and TAB2; Figure 2.12; Wajant and Scheurich, 2011).

The second scenario occurs at late time-points of TNF-R1 stimulation or when NF-κB activation is specifically inhibited (Varfolomeev and Ashkenazi, 2004). Importantly, two different types of Complex II can be distinguished, named Complex IIa and IIb. Complex IIa requires the dissociation of RIP1, TRAF2, and TRADD from the receptor and the subsequent recruitment of FADD and caspase-8 (Micheau and Tschopp, 2003). Differently from Complex IIa, which drives to apoptosis, the Complex IIb is responsible for a distinct type of cell death, named necroptosis, which has recently been object of high interest. Necroptosis is both mechanistically and morphologically distinct from apoptosis. When caspase-8 is activated in Complex IIa, it cleaves and halts the activities of RIP1, RIP3, and the deubiquitinating enzyme CYLD. However, caspase-8 inhibitors (e.g. zVAD) or genetic deletion of caspase-8 or FADD impair RIP1 and RIP3 cleavage and lead to the Complex IIb formation and subsequent necroptosis. Importantly, the activation of necroptosis is tightly controlled by cIAPs, which
are responsible for the proteasome-dependent degradation of active RIP1/3 complexes, and therefore can block the execution of this process (Oberst, 2016).

**Figure 2.12 - Model of TNF-R1 signal transduction** (Ofengeim and Yuan, 2013). Upon ligand binding, TNF-R1 forms the Complex I through the recruitment of several factors, including TRADD, RIP1, TRAF2, cIAP1 and 2. This Complex leads to the recruitment of TAK1 and IKK and drives either to the activation of NF-κB, mediating IκB degradation, or to MAPK activation. Importantly, within 10–15 min after TNF stimulation, the deubiquitylation enzymes disrupt Complex I mediating the deubiquitination of its members. This results in the formation of alternative complexes, named Complex IIa or Complex IIb. Complex IIa includes FADD, caspase-8 and RIP1, and mediates apoptosis. However, the inhibition of caspase-8 leads to Complex IIb, which triggers necroptosis.

### 2.3.2 TNF-R2 complex and its signal transduction

Differently from TNF-R1, TNF-R2 expression is more restricted to specific cell types, including neurons, oligodendrocytes, microglia and astrocytes in the brain (McCoy and Tansey, 2008), endothelial cells, CD4+ and CD8+ T cells (Ware et al., 1991), cardiac myocytes (Irwin et al., 1999), thymocytes and human mesenchymal cells (Bocker et al., 2008). Even
though TNF-R2 lacks DD, it still interacts with TRAF2 and also this receptor can activate the NF-κB pathway in a TNF-R1-independent manner (Borghi et al., 2016). In details, TNF-R2 trimerization allows the recruitment of TRAF2 and consequently of the TRAF2-associated proteins: TRAF1 and cIAPs, thus forming a complex, which interacts with TRAF3 (Figure 2.13). Most of the biological activities triggered by TNF-R2 depend on its interaction with TRAF2, which provides a mechanism for activation of NF-κB, whose translocation to the nucleus promotes the expression of pro-survival target genes, including IAPs and pro-survival BCL-2 proteins (Pahl, 1999). Importantly, inhibition of TRAF2/3 or IAPs reverses the classical NF-κB activation pathway to the non-canonical one (Zarnegar et al., 2008).

![Figure 2.13 - TNF-R2 signal transduction](Cabal-Hierro and Lazo, 2012). TNF-R2 signalling is triggered by the binding with mTNF. This event prompts to the recruitment of TRAF2, which enables the generation of a signalling complex composed by TRAF3, cIAP1 and cIAP2.
2.4 The NF-κB signalling pathway

It is generally accepted that the most important contribution of cIAPs to cell survival and tumourigenesis resides in their ability to regulate NF-κB signal transduction (Zarnegar et al., 2008). Despite its major role in cell survival, NF-κB is also recognised in the modulation of the inflammatory state and innate immunity, and in the control of different steps of cancer initiation and progression (Grilli et al., 1993). The NF-κB family is composed by p65 (RelA), RelB, c-Rel, NF-κB1 and NF-κB2 that associate to form homo- and hetero-dimeric complexes in almost any combination (Figure 2.14; Hayden and Ghosh, 2004). They all share structural features, including a Rel homology domain (RHD; Hoesel and Schmid, 2013), which is required for dimerization, DNA binding, interaction with IκBs, as well as nuclear translocation. In contrast, the transcription activation domain (TA) necessary for the target gene expression is present only in the C-terminus of p65, c-Rel, and RelB subunits (Figure 2.13). Among the 15 potential homo-hetero dimers, three are not able to bind DNA (RelA:RelB, cRel:RelB and RelB:RelB), while other three bind DNA but lack the transcriptional activity (p50:p50, p52:p52, and p50:p52; O'Dea and Hoffmann, 2010). Upon stimulation, NF-κB1 and NF-κB2, which are synthesized as pro-forms (p105 and p100), are proteolytically processed to p50 and p52, respectively. Since neither NF-kB1 nor NF-kB2 contains a TA domain, homo-dimers of p50 and p52 binding to gene promoters act as transcriptional repressors (Chen et al., 2000). However, when p50 or p52 are bound to other NF-κB family members containing a TA domain, such as p65 or RelB, they form a transcriptional activator (Wan and Lenardo, 2009). Different NF-κB subunits are responsible not only for the activation of diverse target genes, but also contain sites for phosphorylations and other post-translational modifications, which enable the cross-talk with other signalling pathways.
Figure 2.14 - NF-κB family members. Adapted from (Hoesel and Schmid, 2013). NF-κB family is composed by five members, namely RelA (p65), RelB, c-Rel, NF-κB1 (p105), and NF-κB2 (p100). All the members contain a RHD, which allows their binding with DNA and the homo- and hetero-dimerization. The C-terminal TA domain is harboured only by the three members: RelA, RelB and c-Rel and mediates their transcriptional activity. The p105 and p100 members show the presence of ankyrin (ANK) repeats motif, which mediates the binding of IκBs to the NF-κB family of proteins and DD motif. The leucin-zipper-like domain (LZ) characterizes only RelB structure.

2.4.1 Activation of the canonical NF-κB pathway

In the canonical pathway, the activation can be mediated by binding of immune receptors including Toll-like receptors (TLRs), Interleukin-1 receptor (IL-1R), TNF-R and antigen receptors with their ligands, such as TNF and lipopolysaccharides (LPS; Ghosh and Hayden, 2008; Perkins, 2007). This interaction drives to the IκBα degradation, with the consequent release of NF-kB and nuclear translocation of p65-containing hetero-dimers, which results in the regulation of target genes (Figure 2.15).
Figure 2.15 - The canonical NF-kB signalling pathway (Ruland, 2011). After TNF binding, the recruitment of TRADD, TRAF2 and cIAP1/2 and RIP1 occurs at the receptor level. This event mediates the activation of IKK complex, which in turn phosphorylates IκBα, thus triggering its degradation. The loss of IκBα allows NF-κB nuclear translocation resulting in the activation of gene transcription. Among the NF-κB target genes there are those that encode for its negative regulators as IκBα.

2.4.2 Non-canonical NF-kB pathway

The molecular mechanisms underlying the non-canonical NF-kB signalling pathway have been investigated only recently. Differently from the canonical one, the activation of the non-canonical NF-kB signalling is slow (within hours) and responds to a different set of stimuli (Razani et al., 2011), e.g. B-cell activation factor (BAFFR), CD40, receptor activator for nuclear factor kappa B (RANK) or lymphotoxin β-receptor (LTβR). Upon stimulation, NIK, the apical inducer of non-canonical NF-kB signalling, phosphorylates and activates IKKα (Figure 2.16), which in turn mediates the phosphorylation of p100 associated with RelB. This event leads to the partial processing of p100 and generation of p52-RelB complex (Sun, 2011). More in-depth, after receptor activation, the complex responsible for NIK degradation,
comprising TRAF3, TRAF2 and cIAP1/2, undergoes cIAP-mediated degradation with the consequent stabilization of NIK. Both TRAFs are responsible for the recruitment of cIAPs, but TRAF3 binds to NIK in a TRAF2-independent manner, while TRAF2 mainly interacts with cIAP1/2.

Figure 2.16 - Non-canonical NF-kB signalling (Cildir et al., 2016). Under resting conditions, NIK is degraded by proteasome in a TRAF2/3-dependent manner. In fact, TRAF2 recruits cIAP1 and 2 and promotes NIK ubiquitination and degradation. Then, after the activation of the non-canonical NF-kB pathway, cIAP1 and 2 mediate TRAF3 degradation therefore resulting in the stabilization of NIK. Once stabilized, NIK phosphorylates IKKα, which in turn phosphorylates p100 leading to its partial degradation to p52. Nuclear translocation of RelB–p52 hetero-dimers allows the expression of target genes.
2.5 IAPs and metastasis

A still less described role of IAPs regards their involvement in the metastatic process. Through the regulation of NF-kB and MAPK signalling pathways, IAPs are able to modulate cell plasticity, motility and invasion (Fulda, 2014c), which are key steps in metastasization. The metastatic process consists of several steps that allow cancer spread from the primary site into the surrounding tissues, and eventually resulting in the colonisation of a secondary site where cells can grow and give secondary tumours (Figure 2.17). During the malignant progression, cancer cells acquire the ability to invade the basal lamina, surrounding tissues and capillaries as single cells (Giampieri et al., 2009) or clumps (Friedl and Gilmour, 2009), extravasate and colonise distant organs whose function is then compromised.

**Figure 2.17 - Cancer cell metastatic process** (Chaffer and Weinberg, 2011). (A) First, the metastatic cascade initiates with cell detachment from the primary tumour. (B) Cancer cells invade the surrounding tissues and intravasate. (C) Once into the blood vessel, cells (CTCs) transit through the circulation to reach the distant organ. (D) At the distant organ, cells extravasate and invade the microenvironment of this secondary site to colonise the new tissue. (E) Cells evade the innate immune response and survive as a single cell or cluster. (F) To give macrometastasis, cancer cells must be able to proliferate and grow.
IAPs have been shown to regulate migration, invasion and metastasis, but the mechanisms underlying these processes are still controversial. On the one hand, several studies support the idea that the targeting of IAPs causes a reduction of cancer cell metastatic potential. These findings would therefore encourage the employment of SMs for the treatment of those types of tumour that are particularly aggressive. This is the case, for example, of triple negative breast cancers (TNBCs). Accordingly, Lopez J et al. (Lopez et al., 2011) showed that cIAP1 deregulated activity can increase cell proliferation and migration, but SM administration is able to prevent this effect. Moreover, they demonstrated that cIAP1 depletion results in vascular tree degeneration *in vivo*, indicating a possible role of cIAP1 in the maintenance of vascular integrity, which is essential for tumour growth and favours the migratory capacity of cancer cells. Nonetheless, it has also been reported that cIAPs regulate Ras related C3 botulinum toxin substrate (Rac1), which is a member of Ras family of small GTPases involved in the control of cell migration and invasion. Notably, XIAP and cIAP1 were shown to interact with Rac1 and lead to its proteasomal degradation (Oberoi-Khanuja et al., 2013). In these settings, IAP depletion mediated by pharmacological or genetic tools would increase Rac1 protein levels, eventually enhancing cell motility and migration. Consistently, Fulda’s group (Tchoghandjian et al., 2013) published the effect of SM BV6 at non-toxic concentrations in glioblastoma cell lines, showing that its administration activates the non-canonical NF-kB signalling and promotes cytoskeleton changes that enhance cell motility.

2.5.1 Role of MAPKs in the metastatic process

The activation of MAPK signalling pathways acts in a variety of cellular responses and mediates several processes such as growth, proliferation, differentiation, migration and
apoptosis (Dhillon et al., 2007). Notably, three distinct MAPK pathways have been characterized: extracellular signal-regulated kinase 1 and 2 (ERK1/2), c-Jun N-terminal kinase/stress-activated protein kinase (JNK/SAPK) and p38 (Figure 2.18).

Figure 2.18 - Schematic representation of the MAPK role in tumours (Reddy et al., 2003). A possible cross-talk exists among the three major MAPK cascades. Despite ERK is mainly responsible for cell invasion and metastasis, it also regulates other cellular processes such as tumour growth and inflammation. At the same times, other pathways besides ERK affect the metastatic process.

Beyond their predominant role in the regulation of the NF-κB signalling pathway, IAPs, and in particular cIAP1 and 2, are also involved in the modulation of the MAPK cascade. Several reports demonstrate that the ubiquitin ligase activity of cIAPs plays a critical role for the efficient activation of MAPK signalling (Gardam et al., 2011; Karin and Gallagher, 2009). Indeed, it has been reported by Varfolomeev and colleagues (Varfolomeev et al., 2012) that the loss of cIAP1 and 2 upon treatment with the IAP antagonist BV6 drastically affects the activation of p38 and JNK.
Among MAPKs, the role of the ERK signalling pathway in the metastatic process has been widely studied. ERK controls a large variety of cellular processes, including proliferation, differentiation, survival, migration, angiogenesis and chromatin remodelling (Dunn et al., 2005; Yoon and Seger, 2006), depending on the cellular settings. Focusing on the metastatic process, it has been shown that sustained ERK signalling enhances the accumulation of genes responsible for angiogenesis, cell migration and invasion. This pathway also favours metastasis formation (Reddy et al., 2003; Giehl, 2005) exerting profound effects on actin cytoskeleton organisation and adhesive structures. Moreover, the metastatic process is favoured by ERK signalling pathway in a double manner: via the apoptosis inhibition and through the increase of tumour cell migration. In the first case, MEK/ERK phosphorylate the pro-apoptotic protein BAD, which dissociates from BCL-2 and allows its anti-apoptotic response. Furthermore, ERK pathway favours the anti-apoptotic MCL-1 protein and inhibits the pro-apoptotic BIM protein, through their phosphorylation (Pachmayr et al., 2017). In the second case, MEK/ERK signalling enhances cell migration by promoting cell movement and contraction through the expression of genes involved in the epithelial mesenchimal transition (EMT) or in the remodelling of tumour microenvironment, such as matrix-degrading proteases (Bae et al., 2013). Therefore, tumour initiation and progression are often caused by the aberrant activation of ERK signalling pathway, which is often caused by the epidermal growth factor receptor (EGFR) hyper-activation.

2.6 EGFR signalling pathway in the metastatic process

EGFR (also named ErbB-1, HER1) is a well-known proto-oncogene, in fact, many reports indicate its crucial role in cancer progression and metastasis. This receptor belongs to
the EGFR family, which comprises other three structurally similar receptors, namely HER2 (ErbB-2), HER3 (ErbB-3), and HER4 (ErbB-4; Wieduwilt and Moasser, 2008). Particular interest has been focused on EGFR since its over-expression and/or hyper-activation characterize about 15–30% breast carcinomas and is associated with poor patient outcomes (Sirkisoon et al., 2016).

2.6.1 EGFR structure and down-stream pathways

EGFR is endowed with a tyrosine kinase activity, which is crucial for the modulation of cellular pathways, which are essential in both normal and cancerous cells. The EGFR activation depends on its binding with specific ligands including epidermal growth factor (EGF), transforming growth factor α (TGFα), amphiregulin or neuregulin (Purba et al., 2017). This event can lead either to the formation of EGFR homo-dimers or hetero-dimers with the other three members of the family (Figure 2.19).

![Figure 2.19 - EGFR signal transduction](image)

**Figure 2.19 - EGFR signal transduction** (Martinelli et al., 2009). EGFR signalling pathway initiates upon ligand binding, which drives the formation of homo-dimer or hetero-dimer with other members of the family. The receptor dimerization causes conformational changes, which induces the EGFR tyrosine kinase activity and
results in the phosphorylation of specific tyrosine residues within the intracellular C-terminal domain. Upon phosphorylation, EGFR acquires the capability to enhance several cellular processes, such as cell proliferation, loss of differentiation, invasion and angiogenesis and block of apoptosis.

The binding of the ligand activates the intracellular tyrosine kinase domain of the dimerized receptor, thus leading to the phosphorylation of specific tyrosine residues within the EGFR intracellular C-terminal domain (Martinelli et al., 2009). Upon phosphorylation, the tyrosine residues provide a docking site for proteins containing Src homology 2 (SH2) domains, such as Grb2, Shc1, p85, PLCγ and JAK1, and trigger the activation of down-stream signalling cascades including the Phosphatidylinositol 4,5-bisphosphate 3-kinase (PI3K/Akt), JAK/STAT, NF-κB, PLCγ/protein kinase-C (PKC), and RAS/MAPK/ERK (Lindsey and Langhans, 2015). These effectors affect several cellular functions such as cell proliferation, loss of differentiation, motility, invasion, angiogenesis and blocking of apoptosis.

Although in normal tissues the presence of EGFR ligands is tightly regulated to ensure the maintenance of cell homeostasis, EGFR is often chronically stimulated in tumour cells. In some cases, the aberrant activation of EGFR, and consequently of its down-stream effectors, is caused by the sustained production of EGFR ligands in the tumour microenvironment. Moreover, in cancer cells, EGFR can be either over-expressed or mutated, causing its constitutive activation. This is the case, for example of head and neck, breast, lung, colorectal, prostate, kidney, pancreas, ovary, brain and bladder tumours (Woodburn, 1999). Remarkably, the correlation between high EGFR expression and poor patient survival renders this receptor a strong prognosis factor in breast, ovarian and head and neck cancers (Fischer-Colbrie et al., 1997; Ishitoya et al., 1989; Sebastian et al., 2006). Not surprisingly, EGFR has emerged as an important target for therapeutic intervention.
2.7 Breast cancer classification

Breast cancers can be classified in five intrinsic subtypes called Luminal A, Luminal B, HER2-enriched, claudin-low and basal-like on the basis of expression levels of hormone estrogen (ER) and progesterone (PR) receptors, HER2, cytokeratins (CKs) 5/6, and claudins 3/4/7 (Prat and Perou, 2011). Tumours positive for ERα and/or PR, but with a low expression of Ki67, are generally defined Luminal A and are responsive to hormone therapy and chemotherapy. Differently, a variable response to chemotherapy has been assessed for Luminal B tumours, which are distinguished from Luminal A tumours due to the higher levels of Ki67. Moreover, the negative expression of ERα and PR characterize both the HER2-enriched tumours and the TNBCs, with HER-enriched tumours expressing high levels of this receptor, while TNBCs are negative. The majority of TNBCs display a basal subtype characterized by strong expression of basal markers such as CK 5, 6 and 17 (Alluri and Newman, 2014).

2.7.1 TNBC: molecular characterization and targeted therapies

TNBCs accounts for about 20 % of the invasive breast cancer cases, characterized by high grade tumours, distant metastasis and low survival. Since TNBCs lack ER, PR and HER2 (Figure 2.20), patients do not benefit from hormonal or trastuzumab-based therapies. Consequently, therapeutic intervention for TNBC women is very limited and patients often show a high risk of recurrence and disease progression. Additionally, TNBC is considered a highly heterogeneous disease and many attempts have been made for its classification. Recently, six molecular subtypes of TNBC have been identified, including basal-like 1, basal-like 2, immunomodulatory, mesenchymal-like (ML), mesenchymal stem–like (MSL), and
luminal androgen receptor, with distinct gene expression profiles and canonical pathways (Lehmann et al., 2011). The high molecular heterogeneity of TNBC disease represents a further barrier in improving survival and in developing targeted therapy for patients (Figure 2.20; Ahn et al., 2016). Currently, TNBC remains the only major type of breast tumour for which U.S. Food and Drug Administration (FDA)- or European Medicines Agency (EMA)-approved targeted therapy is not available, making invasive measures, such as surgery and chemotherapy, the only approaches for the treatment of TNBC patients (Lehmann and Pietenpol, 2015).

2.7.2 Targeting EGFR for anti-cancer treatment

So far, two pharmacological approaches have been proposed to inhibit EGFR activity: neutralizing monoclonal antibodies and small tyrosine kinase inhibitor molecules.

2.7.2.1 Monoclonal antibodies

Among the monoclonal antibodies that have been developed, cetuximab (C225, Erbitux) and panitumumab are currently widely employed in cancer treatment. Cetuximab is
a chimeric IgG1 monoclonal antibody that targets the ligand-binding domain of the EGFR (Figure 2.21). Through its binding with EGFR, cetuximab prevents the receptor activation and subsequent dimerization, and therefore inhibits its signal transduction and hyper-proliferative effects (Ferraro et al., 2013). This drug has been shown to prevent EGFR-dependent primary tumour growth and metastasis. Hence, the employment of cetuximab for clinical use has been approved by FDA for the treatment of patients with wild-type (wt) KRAS, EGFR-expressing metastatic colorectal cancer (mCRC) and recurrent or metastatic head and neck cancers. Activity of cetuximab has been further tested for the treatment of metastatic NSCLC and breast tumours. Panitumumab is a fully human IgG2 targeting the extracellular domains of EGFR. It has been evaluated in clinical trials both in monotherapy and in combination with other agents for the treatment of various cancer types (colorectal and kidney tumours). Along with cetuximab, also panitumumab employment has been approved for treatment of colorectal cancer.

2.7.2.2 Small-molecule EGFR tyrosine kinase inhibitors

Structural studies have identified the ATP-binding site of the intracellular domain of EGFR as responsible for the receptor activity. This observation allowed the development of small-molecule EGFR tyrosine kinase inhibitors (Zhang et al., 2006; Yun et al., 2007). These compounds were designed to compete with the Adenosine 5’ triphosphate and inhibit the intracellular catalytic domain, eventually preventing the EGFR autophosphorylation and its down-stream signalling. These molecules differ in their abilities to bind the ATP-binding pocket—either reversibly or irreversibly—and in their capacities to interfere not only with EGFR but also with other members of the family (Janmaat and Giaccone, 2003). Among the
proposed EGFR inhibitors, gefitinib (Figure 2.21) and erlotinib, have been investigated in preclinical studies and exhibited an encouraging clinical response.

Figure 2.21 - Blocking the EGFR signalling transduction (Graham et al., 2004). After dimerization, EGFR phosphorylated sites residues act as docking sites for several molecules, which cause the activation of diverse signalling pathways. To prevent this event, two strategies have been developed by designing EGFR-specific monoclonal antibodies and small-molecule tyrosine kinase inhibitors.

2.8 EGFR drives metastasis via MAPK signalling pathways

EGFR orchestrates several cellular responses including proliferation, cell motility, angiogenesis, cell survival, and differentiation through the activation of the RAS/MAPK cascade, together with the PI3K/Akt pathway (Schlessinger, 2000). Importantly, many findings provide evidence that MAPKs, and especially ERKs, are involved in cancer initiation and progression (Figure 2.22). In fact, EGFR leads to the activation of the RAS/RAF-MEK1/2-ERK1/2 axis. These signalling pathways are deregulated in approximately one-third of all human cancers and are consequently intensively studied (Dhillon et al., 2007).
Figure 2.22 - EGFR regulation of cellular processes via ERK pathway. Modified from (https://courses.lumenlearning.com/wmopen-biology1/chapter/propagation-of-the-signal/). EGF activates the receptor intrinsic kinase, which in turn induces a large variety of down-stream intracellular signalling pathways. Among these, ERK activation strongly affects cancer behaviour, influencing cell growth, wound healing and tissue repair.

Constitutive activation of ERK signalling, which is found in the vast majority of cancer-associated lesions, is caused by: over-expressed or mutated receptor, sustained autocrine or paracrine production of activating ligands and mutated RAS or BRAF (Figure 2.23). Aberrant activation of the EGFR/MEK/ERK signalling pathway and high levels of ERK1/2 phosphorylation, indicating an elevated activity, have been observed in metastatic sites of breast tumours (Adeyinka et al., 2002). In line with this consideration, MAPK signalling pathway is hyper-activated in TNBCs, where it is associated with both deregulated
proliferation and capability to migrate of malignant cells (Bartholomeusz et al., 2012). Moreover, activation of this pathway has also been linked to higher recurrence rate (Eralp et al., 2008). Finally, emerging evidence links the activation of ERK signalling pathway to the induction of EMT (Xie et al., 2004).

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<th><strong>EGFR overexpression</strong></th>
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<tr>
<td><strong>ERBB2 overexpression</strong></td>
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<td>• Lung adenocarcinoma (35%) (non-small cell)</td>
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<td>• Thyroid: undifferentiated papillary (60%)</td>
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<td></td>
<td>• Melanoma (66%)</td>
</tr>
<tr>
<td></td>
<td>• Colorectal (12%)</td>
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Figure 2.23 - Cancer lesions associated with ERK signalling pathway deregulation (Dhillon et al., 2007).

2.9 EMT process in tumour progression

The essential features of EMT are associated with loss of cell-cell contact and decrease of epithelial features with gain of mesenchymal properties. Several studies have focused on the mechanisms underlying the EMT process supporting its role in tumour cell progression, invasion and metastasis. However, the importance of this process for tumour metastasis is still controversial (Fischer et al., 2015). During the EMT, the down-regulation of E-Cadherin is a crucial step (Acloque et al., 2008) for the reduction of intercellular adhesion junctions and consequent increase of cell detachment. This event is mediated by several E-Cadherin repressors, such as SNAIL, SNAI2 (Bartle et al., 2000), twist family BHLH
transcription factor (TWIST) 1 and 2 (Yang et al., 2004), and the zinc-finger E box-binding homeobox (ZEB) 1 and 2 (Vandewalle et al., 2005). However, it is not clear if these EMT factors function independently or coordinately to activate the program. Notably, the most malignant cancers, including breast cancer, express high levels of Vimentin (mesenchymal marker) along with the decrease of E-Cadherin (epithelial-related), which correlates with poor prognosis (Thiery, 2002). TNBC tumours show high expression of genes associated with EMT (Jang et al., 2015) that are induced by several mediators and, in particular, by the constitutive activation of ERK (Figure 2.24), which sustains cell migration (Huang et al., 2004).

**Figure 2.24 - ERK-mediated regulation of SNAI2 expression.** Adapted from (Kolch et al., 2015). ERK signalling pathway governs EMT by favouring the expression of genes, which are responsible for the regulation of this process, such as SNAIL1 and SNAI2/SLUG. Accordingly, the expression of SNAIL1 and SNAI2/SLUG is accompanied by the decrease of epithelial markers (E-Cadherin), along with the increase of mesenchymal ones (Vimentin).
2.9.1 The ERK signalling pathway promotes tumour aggressiveness

ERK chemical and genetic inhibition results in the reduction of cell migration (Huang et al., 2004). Accordingly, tumour aggressiveness often relies on ERK-mediated phosphorylation of numerous targets, including kinases, phosphatases, transcription factors and cytoskeletal proteins (Dhillon et al., 2007). Notably, several mediators of the EMT process are found among these targets, e.g. SNAIL1, SNAI2 (Chang et al., 2011), TWIST1 and ZEB1, which are commonly high expressed in TNBCs (Figure 2.25).

![Figure 2.25 - EMT features and molecular portrait](image)

**Figure 2.25 - EMT features and molecular portrait** (Shih and Yang, 2011). EMT is a crucial step for tumour progression and invasion and promotes metastasis formation, because of the increased cell motility that follows the acquisition of mesenchymal features. Furthermore, EMT favours apoptosis resistance to chemotherapeutic drugs and the expression of stem cell-like features.

2.9.2 The role of SNAI2 in cancer metastasis

The expression of the transcription factor SNAI2 is sustained by EGFR down-stream pathways, including PI3K-Akt and RAS-MAPK. SNAI2 belongs to the highly conserved
Snail/Scratch superfamily and it is characterized by two domains: the highly conserved SNAG (Snail/Gfi) domain in the N-terminal region and the C2H2 type zinc fingers in its C-terminal region (Nieto, 2002) Figure 2.26).

![Figure 2.26 - Structural domains of SNAI2. Modified from (Shih and Yang, 2011). The zinc finger transcriptional factor SNAI2 contains an N-terminal SNAG domain and C-terminal zinc finger domains. Both of these domains are responsible for SNAI2 repressor activity. The NES domain allows the nuclear export signal.](image)

Several agents, including EGF, fibroblast growth factor (FGF), hepatocyte growth factor (HGF), transforming growth factor β (TGFβ), bone morphogenetic proteins (BMPs), WNTs and Notch (Figure 2.27), activate SNAI2 expression. Among these factors, EGF is important for SNAI2 high expression in TNBCs, which enhances cell motility, metastatic potential (Bailey et al., 2012; Phillips and Kuperwasser, 2014) and resistance to detachment-induced cell death.
Figure 2.27 - Pathways responsible for SNAI2 expression (Barrallo-Gimeno and Nieto, 2005). The transcription factor SNAI2 is controlled by several pathways. Its modulation occurs not only at the transcriptional levels, but also through the regulation of its subcellular localization.

Therefore, the association of SNAI2 with cancer aggressiveness and resistance to therapy renders this protein an attractive therapeutic target in breast cancers (Figure 2.28). However, only few compounds have been designed to directly target SNAI2 (Harney et al., 2009) and some indirect approaches have been developed to inhibit its expression and/or activity (Ferrari-Amorotti et al., 2014; Ferrari-Amorotti et al., 2013). In this way, the invasive and migratory behaviours of cancer cells have been attenuated.
Figure 2.28 - Role of SNAI2 in tumour metastasis. Adapted from (Chimge and Frenkel, 2013). SNAI2 controls both early metastatic events including EMT, invasion, intravasation and late stages, thus favouring cancer cell colonisation of secondary sites.

Beyond EMT regulation, SNAI2 increases cancer aggressiveness favouring cancer cell stem-like features (Luanpitpong et al., 2016). In fact, a correlation exists between EMT and cancer stem-like cell (CSC) enrichment, and in line with this findings, it has been reported that SNAI2 controls the levels of SOX2 (Samanta et al., 2016) and SOX9 (Luanpitpong et al., 2016). These proteins belong to the SOX (Sry-related HMG Box) family, which is involved in the maintenance of self-renewal and pluripotency in embryonic stem cells, as well as in adult tissue progenitors (Sarkar and Hochedlinger, 2013). Therefore, SNAI2 plays a pivotal role in the regulation of cell motility, survival and cancer cell “stemness” and contributes to the dissemination of cancer cells from the primary tumours to the secondary sites. In light of these observations, it would be important to understand the molecular mechanisms of
action and regulation of SNAI2 in order to design novel approaches to interfere with its expression and contrast the metastatic process.

2.10 EGFR intracellular trafficking

So far, the mechanisms leading to EGFR over-expression in TNBCs are not completely clear. Increased levels of this receptor have been linked to the presence of mutated BRCA (Nakai et al., 2016), but also other aspects contribute to EGFR levels. For example, it has been shown that a reduced degradation can sustain its levels (Zhang et al., 2013). EGFR down-regulation occurs at the end of an intricate process resulting in signal attenuation, which derives from the removal of the receptor from the cell surface (Peschard and Park, 2003). Upon ligand-binding, the major mechanism of EGFR internalization is clathrin-mediated endocytosis (CME), through which the receptor is internalized by clathrin-coated pits and then directed to early endosomes where the cargo is delivered after fusion. Following the delivery to the early endosome, the receptor can be either recycled to the cell surface or sorted to late endosomes and lysosomes for being degraded. An alternative pathway for EGFR internalization is clathrin-independent and it is therefore called clathrin-independent endocytosis (CIE; Sigismund et al., 2005). It has been published that high concentrations of EGF drives to CIE rather than CME (Figure 2.29).
Figure 2.29 - EGR endocytosis and endosomal sorting (Haglund and Dikic, 2012). Upon EGF binding, the receptor undergoes endocytosis, which can occur in a clathrin-independent (left) or clathrin-mediated (right) manner. In both cases, after being internalized, the receptor is routed to early endosomes and then sorted to late endosomes (MVE) for lysosomal degradation (ubiquitylated receptors) or recycled to the plasma membrane.

2.11 EGFR ubiquitination is a crucial step for its degradation

In addition to controlling localization, function and stability of target proteins, ubiquitination appears as a crucial event in EGFR endocytosis and degradation. Ubiquitinated EGFR is mainly degraded through lysosomal vesicles (Futter et al., 1996; Marmor and Yarden, 2004) and not by “classical” proteasomal degradation.

2.11.1 EGFR degradation induced by c-CBL

Commonly known as being a negative regulator of various activated receptor tyrosine kinases (RTKs) such as EGFR and c-MET, c-CBL plays a crucial role in EGFR endocytic
trafficking. Notably, c-CBL is a member of the evolutionarily conserved Cbl family of cytoplasmic proteins and because of its crucial role in the attenuation of receptor signalling, c-CBL deregulation can lead to malignant diseases. Accordingly, Zhang J. et al. (Zhang et al., 2013) reported that c-CBL inhibition correlates with poor prognosis in glioblastoma patients. Interestingly, it was shown that if c-CBL is unable to correctly regulate EGFR, this receptor does not undergo lysosomal degradation. This block eventually results in extended EGFR-signalling activities (Figure 2.30).

![Figure 2.30 - c-CBL-mediated ubiquitination leads to EGFR endocytosis and degradation. Adapted from (Zhang et al., 2013). c-CBL crucially mediates EGFR ubiquitination and degradation. The impairment of c-CBL activity results in enhanced signalling which contributes to cancer progression.](image)

Upon ligand-binding, EGFR undergoes dimerization and autophosphorylation of several tyrosine residues in the cytoplasmic tail. These phosphorylations represent the prerequisite for clathrin-dependent endocytosis. In particular, phosphorylation of tyrosine 1045 (Tyr-1045) is crucial for the direct binding of c-CBL to EGFR (Levkowitz et al., 1999). It has been recently reported that the hypophosphorylation at Tyr1045 allows EGFRvIII to
evade degradation (Grandal et al., 2007). In this setting, c-CBL is not recruited to EGFR, which therefore it is not ubiquitinated correctly (Figure 2.31).

Figure 2.31 - Phosphorylation of Tyr-1045 is crucial for c-CBL-mediated EGFR degradation (Grandal et al., 2007). The binding of c-CBL requires the phosphorylation of EGFR at Tyr-1045. Consequently, the mutant form of EGFR, named EGFRvIII, which is hypophosphorylated at Tyr-1045, avoids the ubiquitination by c-CBL and is less degraded.

2.11.2 RAB family proteins regulate the EGFR endocytic trafficking

Once EGFR is in early endosomes, it can be routed to the late endosome and lysosome for degradation, or recycled to the plasma membrane. If the receptor is targeted for degradation, it is first internalized in multivesicular body (MVB; Stahl and Barbieri, 2002), and subsequently released to the lysosomes. RAB proteins have been identified as key
regulators of EGFR endocytic trafficking and mediate this process. The RAB family consists of small GTPases proteins (Somsel Rodman and Wandinger-Ness, 2000) and is composed by over 60 members that are characterized by similar structure and properties, but exert different functions in the regulation of intracellular vesicles trafficking and fusion reactions. So far, only a few RAB proteins have been described as being regulators of EGFR endocytic trafficking (Ceresa, 2006) namely RAB5, RAB11 and RAB7. RAB5 and 11 are established markers of early (Barbieri et al., 2004; Konstantinopoulos et al., 2007) and recycling endosomes (Cullis et al., 2002), respectively. Conversely, RAB7 is a marker of late endosomes, and therefore drives EGFR to degradation (Ceresa, 2006; Figure 2.32).

Figure 2.32 - Different RAB family members are responsible for EGFR recycling and degradation (Francavilla et al., 2016). Once in early endosomes, EGFR can be recycled to the cell surface or further routed to late endosomes and lysosomes for degradation. While RAB11 is a marker of the recycling process, RAB7 is a marker of late endosomes and co-localizes with degrading EGFR.
2.12 The role of LRIG1 in EGFR down-regulation

Together with c-CBL, the leucine rich repeat and immunoglobulin-like domain protein-1 (LRIG1) has recently been identified as a negative regulator of c-MET and the ErbB family members (Figure 2.33). Particularly, Gur et al. reported the involvement of LRIG1 in increasing EGFR ubiquitination and degradation, which occurs by enhancing the recruitment of c-CBL to the receptor (Gur et al., 2004). However, while LRIG1 mediates EGFR degradation in a c-CBL dependent manner (Gur et al., 2004), c-MET degradation does not require ubiquitination and is mediated by LRIG1 in a c-CBL-independent manner (Shattuck et al., 2007).

![Figure 2.33 - The role of LRIG1 in EGFR degradation](image)

*Figure 2.33 - The role of LRIG1 in EGFR degradation* (Wang et al., 2013). LRIG1 negatively regulates EGFR via enhancing its interaction with c-CBL. The presence of LRIG1, in fact, favours EGFR degradation by accelerating the recruitment of the E3 ligase c-CBL through a CBL-binding domain in the LRIG1 cytoplasmic tail. This mechanism enhances EGFR ubiquitination and lysosomal degradation.
LRIG1 has been described as a tumour suppressor due to its capability to inhibit cell proliferation and cancer growth (Hedman et al., 2002) and its expression has been analysed in many human cancers. Importantly, gene expression analysis of five cancers, including breast, lung, bladder, glioma and melanoma (Rouam et al., 2010), revealed that LRIG1 decreased expression correlates with poor survival. Being a transcriptional target of estrogen receptor-alpha (ERα), LRIG1 expression is high in Luminal A (ER-positive) breast cancers and low in the basal-like subtype (Wang et al., 2013). Interestingly, LRIG1 low expression contributes to the aggressiveness of basal-like tumours. In fact, a negative correlation has been found between its expression and the presence of mesenchymal markers (Yokdang et al., 2016).

2.13 EGFR and c-MET cross-talk

Together with EGFR, also the activation of c-MET mediated by its ligand, the hepatocyte growth factor (HGF), is known to promote the survival of many cell types. Furthermore, c-MET enhances tumour invasion via increased ERK phosphorylation, and this results in a high metastatic potential of cancer cells. Importantly, the interaction between EGFR and c-MET has been found in tumour but not in normal cells (Jo et al., 2000). This evidence suggests that a tumour-specific cross-talk exists between EGFR and c-MET receptors and promotes the activation of their shared down-stream pathways (Figure 2.34). Many signalling pathways are in common between both receptors and therefore several studies support a possible synergism between EGFR and c-MET, which favours tumour growth and aggressiveness. However, the underlying mechanisms responsible for this functional interaction remain unknown. Furthermore, Engelman and colleagues reported
that patients affected by NSCLC are initially sensitive to treatment with the EGFR inhibitor gefitinib, but later become resistant because of c-MET gene amplification. Emerging evidence shows that about 20% of acquired resistance to EGFR-targeted therapies derives from the amplification of the c-MET receptor (Corso and Giordano, 2013). Accordingly, breast cancer cells acquire resistance to EGFR TKIs after HGF exposure and the HGF-induced cell survival results abrogated in EGFR-depleted cells, indicating a critical role for EGFR/c-MET cross-talk (Mueller et al., 2012). Intriguingly, this information supports the employment of simultaneous treatment with anti-EGFR and anti-c-MET drugs to restore the sensitivity and efficiency of tumour treatment (Yu et al., 2013).

![Functional interaction between EGFR and c-MET](image.png)

**Figure 2.34 - Functional interaction between EGFR and c-MET** (Zhang et al., 2015). A cross-talk between EGFR and c-MET down-stream signalling pathways occurs thereby enhancing tumour malignant phenotype. The two receptors elicit similar signal transduction pathways and therefore their synergism increases the strength and duration of shared signalling pathways. EGFR and c-MET are often co-expressed in tumour with a high grade of aggressiveness and their combined activity promotes cell survival, proliferation and migration.
3 AIM OF THE STUDY

Despite the progress in the understanding of the molecular and cellular basis driving to metastasis formation, this process remains the major cause of mortality. The central purpose of this study is therefore to characterize the role of IAPs in the metastatic cascade.

The specific aims of this work are:

- Investigate the effect of SM83-mediated depletion of IAPs on metastasis formation in vivo
- Determine the mechanism of SM83-induced down-regulation of SNAI2
- Study the role of IAPs, and in particular of cIAP1, in regulating EGFR-mediated SNAI2 expression
- Investigate the role of cIAP1 in regulating EGFR levels.
4 MATERIALS AND METHODS

4.1 Reagents

4.1.1 Buffers and Solutions

Freezing solution

- 60 % RPMI or DMEM supplemented with
  - 1 % L-Glutamine (LONZA)
  - 1 % NEAA (LONZA)
  - 1 % Sodium pyruvate (LONZA)
  - 1 % Hepes (LONZA)

- 30 % FBS

- 10 % DMSO (Sigma-Aldrich)

SDS lysis buffer

- 5 % SDS

- 125 mM Tris HCl pH 6,8

- Milli-Q water up to volume

Mix of proteases and phosphatases inhibitors

- 200 µg/ml Aprotinin

- 200 µg/ml Leupeptin

- 200 mM EDTA

- 200 mM β-glycerophosphate
- 200 mM sodium fluoride
- 200 mM sodium orthovanadate
- 50 mM sodium pyrophosphate
- Milli-Q water up to volume

**Protein loading buffer 10X (ZAP)**

- 0.25 M Tris HCl pH 6.8
- 30 % glycerol
- 8 % SDS
- 0.02 % BFB
- 10 % β-mercaptoethanol
- Milli-Q water up to volume

**Running Buffers**

MES and MOPS were purchased from Thermo Fisher Scientific

**Transfer buffer**

- NuPAGE Transfer Buffer 1X (Thermo Fisher Scientific)
- 20 % methanol
- Milli-Q water up to volume

**PBS-T**

- PBS (LONZA, Cat. #17-516F)
- 0.1 % TWEEN 20
**Blocking buffer**

- 4% non-fat dry milk (Blotting-Grade Blocker, BIO-RAD)
- 0.01% Tween-20 (Sigma-Aldrich)
- 1X PBS

**ELB buffer**

- 150 mM NaCl
- 50 mM Hepes
- 5 mM EDTA
- 0.5% NP40
- Milli-Q water up to volume
- pH 7.5

**Nucleic acid loading dye 6X**

- 0.25% bromophenol blue powder
- 30% glycerol
- Milli-Q water up to volume

**TAE buffer 1X**

- 20 mM Tris acetate
- 10 mM EDTA
- 2.85% acetic acid
- Milli-Q water up to volume
**Antibody solution**

- 5 % Albumin from Bovine Serum (BSA; Sigma-Aldrich),
- 0.01 % Tween-20,
- 0.01 % Sodium Azide purchased from Sigma-Aldrich,
- 1X PBS

**4.2 Cell cultures**

The human breast adenocarcinoma MDA-MB231, BT549, MDA-MB157, SkBr3 and HCC1937 cell lines were cultured in Roswell Park Memorial Institute (RPMI)-1640 medium (LONZA Group, Basel, CH) supplemented with 10 % fetal bovine serum (FBS; EuroClone, Milan, IT), 2 mM L-glutamine, sodium pyruvate, non essential amino acids (NEAA) and penicillin (100 U/ml)/streptomycin (100 μg/ml; all from LONZA). SUM149 and SUM159, together with the human hTERT-mammary immortalized epithelial HME and MCF10A cell lines- parental and bearing the EGFR delE746A750 mutation (HME EGFR and MCF10A EGFR hereafter, respectively) -were cultured in Dulbecco’s Modified Eagle’s medium-F12 (DMEM-F12; Gibco), supplemented with 10 % FBS, 2 mM L-glutamine, 20 ng/ml EGF (Cat. #GRF-10544, Selleck Chemicals, Munich, D), 10 μg/ml insulin (Sigma-Aldrich, St. Louis, MO, USA), 500 μg/ml hydrocortisone (Sigma-Aldrich). BT474, MDA-MB453, T47D and MDA-MB361 cell lines were maintained in DMEM (Gibco-Thermo Fisher Scientific, Massachusetts, USA) supplemented with 10 % FBS and 2 mM L-glutamine. HEK293FT (Thermo Fisher Scientific) highly transfectable cell line were cultured in DMEM with 10 % FBS and used for lentiviral production. All cells were grown at 37 °C in fully humidified atmosphere with 5 % CO2.
MDA-MB231 cell lines were purchased from American Type Culture Collection while all the other breast cancer cell lines employed for this thesis were kindly provided by Dr Elda Tagliabue. The mouse 4T1 mouse mammary carcinoma were a kind gift from Dr Mario Colombo, while human mammary immortalized epithelial HME and MCF10A cell lines were all kindly provided by Prof. Alberto Bardelli. All cell lines have been checked for their identity (STR characterization) and were mycoplasma-free as determined by Takara Mycoplasma Detection Kit (Clontech, Mountain View, CA, USA).

4.3 Cell viability assay

Cell viability was established using the CellTiter-Glo (Promega, Madison, USA). According to the manufacturer’s instructions, cells were seeded in white optical 96-well plates and treated as indicated. Medium was discarded at the end of the experiments and replaced with 60 µl of reagent diluted 1:4 in phosphate-buffered saline PBS. After 15 min of shaking at RT, luminescence was measured with a Tecan Ultra plate reader. Viability of cells silenced with indicated siRNAs (Table 4.1) was shown as a percentage compared to untreated or mock treated cells.

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</table>

Table 4.1 - List of Dharmacon siRNA pools (GE Healthcare, Lafayette, CO, USA).

4.4 Cell treatments

To deplete cIAPs, cells were treated with 100 nM SM83 whose synthesis has recently been described (Manzoni et al., 2012). The activation of TNF-Rs was achieved by the administration of recombinant TNF (Emmerich et al., 2011) produced by the University of Milan using a plasmid encoding for tagged human TNF kindly provided by Prof. Henning Walczak. To specifically activate the non-canonical NF-κB pathway, the recombinant human TWEAK provided by Prof. Harald Wajant (Salzmann et al., 2013) was employed. In combination therapy with SM83 the necroptosis inhibitor Necrostatin-1 (Enzo Life Sciences, Plymouth Meeting, PA, USA) and Pan-caspase inhibitor z-VAD(OMe)-FMK (BIOMOL) were used. Cells were treated with the PI3K and AKT (LY294002 and Triciribine, Enzo Life Sciences), MEK (U0126, Calbiochem, Merck KGaA, Darmstadt, D) and p38 (SB203580, Selleck Chemicals) inhibitors. EGFR activation was induced through the administration of 20 ng/ml EGF (Cat. #GRF-10544, Selleck Chemicals) or TGFα (#100-16A, Peprotech, London, UK) in cells serum-starved overnight using medium supplemented with 0.1 % FBS. Cetuximab employed to specifically block EGFR was provided by the pharmacy of the Fondazione IRCCS Istituto Nazionale dei Tumori (Milan, IT). When needed, protein levels were stabilised with the lysosomal inhibitor chloroquine (CLQ, #C6628) from Sigma-Aldrich, used at 100 μM and added to the cell growth medium 1 h before EGFR stimulation. The protein synthesis
inhibitor cycloheximide (CHX, #239764, Calbiochem) was added to BT549 cells at 100 μM 30 min before treatments, while half of the dose was employed in MCF10A cells.

4.5 In vivo experiments

Experiments were approved by the Ethics Committee for Animal Experimentation of the Fondazione IRCCS Istituto Nazionale dei Tumori (INT) of Milan according to institutional guidelines and by the Italian Minister of Health (Projects INT_12_2011 and INT_02_2015). Mice were maintained in laminar flow rooms keeping temperature and humidity constant. Mice had free access to food and water and were weighted twice a week.

In vivo effect of SM83 was assessed using the breast cancer xenograft models, non-obese diabetic/severe combined immunodeficiency disease (NOD/SCID) mice. Animals were engrafted in the left flank by subcutaneous (sc) injection of 200 μl physiological saline containing 5x10^6 MDA-MB231. Randomization occurred at the day 13, when mice were treated with ip or iv injections with 5 mg/kg SM83, 5 times/week to a total of 15 injections. Tumour growth was evaluated by biweekly measurements of tumour diameter with a Vernier caliper and tumour volume (TV) was calculated according to the formula: 4/3 x 3.14 x (l/2) x (w/2) x (h/2) where l, w and h are length, width and height, respectively. Biochemical analysis and gene expression analysis of the primary tumour were performed in mice sacrificed 6 h after the last injection. In metastasis studies, NOD/SCID mice were killed 2 weeks after the last SM83 administration and, together with subcutaneous nodules, lungs were collected and formalin-fixed/paraffin-embedded for IHC detection of metastasis with anti-human Vimentin antibody (M0725, DAKO-Agilent, Santa Clara, CA, USA).
The anti-metastasis effect of SM83 was also tested using immuno-competent BALB/c mice injected with $1 \times 10^4$ 4T1(cl5) cells on day 0 into the mammary fat pad. After 13 days, mice were treated with ip injections of SM83 at a dose of 5 mg/kg body weight, three times/weeks for 2 weeks. The progression of metastasis was assessed 2 days after the last SM83 injection.

4.6 Gene expression profiling and bioinformatics

MDA-MB231 nodules were collected from NOD/SCID xenografts 6 h after the last injection of SM83 and cut in pieces. One piece was lysed for RNA extraction using RNeasy Mini Kit (Qiagen, Hilden, Germany) and then total RNA was retro-transcribed using SuperScript II Reverse Transcriptase kit (Thermo Fisher Scientific) for gene expression profiling (GEP). GEP was performed by the Functional Genomics and Bioinformatic Core Facility of INT using Illumina HumanHT-12 v4 Expression BeadChip arrays (Illumina, San Diego, CA, USA).

Raw data were pre-processed using the R/Bioconductor package “lumi” (Du et al., 2008) that provides statistical methods for analysis of Illumina microarray data. Data were Log2-transformed and normalized using the robust spline normalization method. Probes not annotated to HUGO gene symbols were filtered out. For the remaining probes only those with a detection p-value < 0.01 (a measure of the confidence that a probe is expressed above the background level, defined by negative control probes) in at least one sample were considered. Finally, multiple probes mapping to the same gene were collapsed selecting the probe detected in the highest number of samples.
Differential expression analysis was performed using the “limma” package (Phipson et al., 2016) that combines linear models with moderated t-statistic to identify differentially expressed genes across experimental conditions. The moderated t-statistics has the same interpretation as an ordinary t-statistic except that the standard errors are moderated across genes, i.e., squeezed towards a common value, using a simple Bayesian model. P-values obtained from limma were adjusted for multiple-testing using the Benjamini-Hochberg false discovery rate (FDR) to reduce the number of false positives. Genes showing an absolute fold change ≥ 1.5 and an FDR < 0.05 were considered significantly differentially expressed. Expression profiles are deposited in the Gene Expression Omnibus (GEO) repository with accession number GSE98691.

For public gene expression datasets, normalized data for breast cancer cell lines were downloaded from ArrayExpress repository (https://www.ebi.ac.uk/arrayexpress/) with accession number E-MTAB-181 (Heiser et al., 2012). RNA-Seq level-3 expression data for TCGA breast cancer patients were downloaded from Firehose portal (http://gdac.broadinstitute.org/) with accession date 2016-01-28. Data were normalized using the trigger mean of M-value method (Robinson and Oshlack, 2010) and transformed in logarithmic scale (base 2). Correlation between continuous variables was calculated using the Pearson’s correlation coefficient.

4.7 Wound healing-based migration assay

Wound healing experiments were performed using Culture-Insert in µ-Dish 35 mm (Ibidi). To assess cell migration, 4x10^4 MDA-MB231 cells were reverse transfected in the culture inserts in 12-well plates. After 72 h, inserts were removed, cells were washed once to
eliminate detached cells and multi-well plates were put in a Cell-IQ instrument (CM-
Technologies) to test cell motility. Images were taken every hour for 24 h and analysed with
the provided software to measure the gap area.

Wound healing assay was performed using the same protocol described above, also
to compare the migration capability of MDA-MB231, BT549, HCC1937 and T47D cell lines and
then to assess the effect of cIAP1 depletion on cell motility in BT549 and MCF10A, either wt
or bearing mutated EGFR, stably silenced for cIAP1 compared to control. Data obtained by
Cell-IQ instrument were analysed and used to draw a sigmoidal curve with GraphPad Prism
thus determining the time necessary to close half of the wound area.

4.8 Gene knock-down by silencing (reverse protocol)

To achieve transient knock-down of target genes, cells were transfected with
indicated short interfering RNAs (siRNAs, Table 4.2) using a reverse transfection protocol in
which siRNAs (Qiagen) and RNAiMAX (Thermo Fisher Scientific) have been employed. A mix
containing 3.25 µl of RNAiMAX in 100 µl Optimem (Gibco) and another one with 3.25 µl of
siRNA (20 µM) stock were prepared, and after 5 min at RT were combined and left at RT for
30-40 min. In the meantime, cells were trypsinized, counted and about 0.25 x 10^6 cells were
cultured in a 6-well plate in a final volume of 2 ml medium without antibiotics. After the
incubation, the siRNA/RNAiMAX mix was added on top of the cells in the culturing well. Cells
were incubated for 72 h to have an efficient knock-down in all cases. If cells needed to be
stimulated, 48 h after transfection cells were serum-starved for the following 24 h. In each
experiment, scramble siRNAs (siCtr) were used as control and siRNA targeting an essential
gene (ubiquitin, UBB) served to evaluate the transfection efficiency. For transfection in 96-
well plates, mixes were prepared with 0.25 µl siRNAs plus 0.25 µl RNAiMAX and 10^4 cells were seeded in 100 µl final volume.

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<td>Hs_MET_6</td>
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<td>Dharmacon</td>
</tr>
<tr>
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<td>siGENOME SMARTpool for SNAI2</td>
<td>M-017386-00</td>
<td>Dharmacon</td>
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</tbody>
</table>

Table 4.2 - List of siRNAs used in this thesis.

4.9 Lentiviral transduction for stably knock-down

MDA-MB231 cells were engineered to stably knock-down SNAI2 through lentiviral-mediated gene delivery. Lentiviral particles containing the pGFP-C-shLenti SNAI2 (NM_003068) were purchased from OriGene. Target cells (5×10^4 cells) were cultured with medium containing lentiviral particles for 48 h before addition of puromycin (0.5 µg/ml) for selection. After selection, transfection was confirmed by protein expression analyses.
To stably knock-down cIAP1, lentivirus was produced by transfection of HEK293FT packaging cells with the pLKO.1-cIAP1 shRNA (Cat. #44129, Addgene). Lentiviral particles were collected 48 h after transfection and filtered through a 0.45 μm filter.

4.10 Gene over-expression

To ectopically express EGFR and c-CBL, lentiviral plasmids were used to produce lentiviral particles in HEK293FT packaging cells. Viruses were obtained by reverse transfection of HEK293FT cells following manufacturer’s protocol. After 24 h, medium was replaced with fresh medium that was collected after further 24 h and used to transduce BT549 and MCF10A cells. The plasmid for human EGFR (Myc/Flag-tagged) ectopic expression was purchased by Origene Company (#RC217384L1; Rockville, MD, USA), while the c-CBL-expressing vector (RRL-CMV-CBL) was kindly provided by Prof. Pier Paolo Di Fiore.

To over-express wt and mutated cIAP1, cells were seeded the day before transfection, using a medium without antibiotics, in order to have 80-90 % of confluent cells the following day. BT549 cells were transfected with pcdna3.1 plasmids kindly provided by Jon Ashwell (Addgene plasmid #8311 and #8337, respectively), according to the manufacturer’s instructions. Briefly, DNA was complexed with Opti-MEM, as well as Lipofectamine 2000 (ratio DNA : Lipofectamine = 1 : 2). Mixtures were incubated 5 min at RT and then combined to allow the formation of the liposomes-DNA complexes; 30 min later, the solution was added to the cells and the desired protein expression was verified 48 h later.
4.11 Western blot analysis

4.11.1 Preparation of total cell extracts

Cells were trypsinized and harvested by centrifugation at 4500 rpm for 5 min at 4 °C. After washing with PBS supplemented with 0.1 mM Na$_3$VO$_4$ to inhibit phosphatase, cells were lysed by boiling in 60-100 μl SDS lysis buffer supplemented with the mix of phosphatases and proteases inhibitors. Samples were then sonicated with 20 % amplitude for 20 sec (Branson Digital Sonifier) to dissolve DNA molecules, and then centrifuged at 13000 rpm for 20 min at room temperature (RT). Cleared supernatants were transferred to a new tube and frozen at -20 °C.

4.11.2 Quantification of total cell extracts

Bicinchonic acid (BCA)-containing protein assay was used to determine the protein content of total cell extracts, according to the manufacturer’s instructions (QuantumMicro Protein, EuroClone). Briefly, 2 μl of lysate were incubated with 148 μl water and serial dilutions of bovine serum albumin (BSA) used as standard protein, in a 96-well plate. BCA solution was added in a 1:1 ratio. After incubation at RT, absorbance was measured at 485 nm using Ultra microplate reader (Tecan). Protein concentration was determined by interpolation with the curve obtained with the standard BSA.

4.11.3 SDS-PAGE

Proteins were separated according to their molecular weight using pre-cast 4-12 % Bis-Tris NuPAGE gels (Thermo Fisher Scientific). Cell lysates were mixed with 4 x reducing SDS-Sample buffer and heated for 10 min at 99 °C. Before loading, protein samples were
prepared by adding ZAP solution to 20-50 μg of proteins and samples were denatured for 10 min at 99 °C. As a molecular weight standard, Page Ruler Plus Pre-Stained Protein Ladder (EuroClone) was used. The electrophoretic separation was achieved by applying a constant voltage in MES or MOPS buffer. As suggested by the manufacturer, 500 μl of NuPAGE antioxidant were added to the chamber to protect reduced disulfide bonds and sensitive amino acids from oxidation, thus allowing proper protein migration in reducing conditions. Proteins within the gels were then blotted onto PVDF Immobilon-P Transfer Membrane (Millipore), previously activated with 100 % methanol, rehydrated in Milli-Q water, and equilibrated in transfer buffer. Transfer was carried out using the XCell II blot module (Thermo Fisher Scientific). Transfer sandwich was composed of three sponges; three 3MM wetted papers (Whatman), the gel, PVDF and three additional 3MM wetted papers. The sandwich was put into the XCell SureLock Mini-Cell, the blot module was filled with transfer buffer and the outside chamber with distilled water; transfer of proteins was carried out at 180 mA for 2 h. Membranes were incubated with blocking buffer (4% non-fat milk dissolved in PBS plus tween 0.1 %, PBS-T) for 30 min and then incubated overnight with the indicated primary antibodies. Membranes were then washed 3 times in PBS-T and incubated 1 h with the appropriate horseradish peroxidase-conjugated secondary antibody (Sigma). After washing in PBS-T, proteins were detected by electrochemiluminescence (ECL) reaction, by exposure of films to the membranes after incubation with luminol-based chemiluminescent substrates (Pierce).
4.12 Immunoprecipitation

Immunoprecipitation (IP) was performed in BT549 cells over-expressing FLAG-tagged EGFR for the identification of EGFR binding proteins (co-IP). To this end, cells were collected, washed 1 time with PBS supplemented with Na$_3$VO$_4$ and lysed in ice-cold ELB buffer supplemented with the mix of proteases and phosphatases inhibitors (hereafter ELB+), at 4 °C with rotation for 30 min. Samples were then centrifuged at 13000 rpm for 15 min at 4 °C, supernatant was transferred in a new low-retention microcentrifuge tube and 1 mg of total protein extract was incubated at 4°C for 3 h with 30 µl of the 50 % slurry of anti-FLAG M2 Affinity Gel (#A2220, Sigma-Aldrich) previously washed with ELB+ buffer three times. After the incubation time, resins were resuspended in 15 µl of SDS lysis buffer and 5 µl ZAP solution, denatured for 10 min at 99 °C and stored at −20 °C. The bound polypeptides were analysed by SDS-PAGE and immunoblotting with indicated antibodies (Table 4.3).

<table>
<thead>
<tr>
<th>Target antigen</th>
<th>Cod.</th>
<th>Immunoglobulin type</th>
<th>Host organism</th>
<th>Dilution</th>
<th>Source</th>
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<td>cIAP2</td>
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<td>Cell Signaling</td>
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</table>

Table 4.3 - List of antibodies employed in this thesis.

4.13 Stripping of western blot membranes

Western blot membranes were incubated with stripping buffer at 37 °C for 30 min and then washed for 3 times with PBS-T, followed by incubation in blocking solution and a new round of probing.

4.14 Immunofluorescence

BT549 cells were trypsinized, resuspended in 3 ml of cold RPMI with 10 % FBS and spotted on glass microscope slides using cytospin (Cytospin 2, Shandon). According to Cytospin protocol, slides and filters were placed into appropriate slots in the cytospin with the cardboard filters facing the center of the cytospin. About 100-200 μl of cell suspension were put into each of the wells and spinned at 500 rpm for 5 min. Then slides were dried
overnight fixed with 4 % paraformaldehyde for 10 min, permeabilized for 10 min at RT with 0.2 % Triton X-100 and blocked in PBS, 3 % BSA, 0.1 % Tween-20. Cover slips were then incubated overnight at 4 °C in a humidified chamber with the primary antibody diluted in PBS 2 % BSA overnight. After three washes in PBS, the appropriate secondary fluorescent Alexa Fluor conjugated antibody (Thermo Fisher Scientific), diluted in PBS 2 % BSA was added to the cells for 1 h and the plate was kept in the dark. After incubation, three washes with PBS were performed and nuclei were counterstained with DAPI (0.5 µg/ml in PBS) for 10 min. Then coverslips were mounted on glass microscope slides with ProLong Gold reagent mounting solution (Thermo Fisher Scientific). Fluorescence images were acquired using a fluorescence microscopy and digital image acquisition on a Nikon Eclipse E1000 equipped with a DSU3 CCD camera.

4.15 In situ Proximity Ligation Assay (PLA)

Duo Link in Situ reagents (Duolink In Situ PLA; Sigma-Aldrich) were employed in this thesis following the manufacturer’s guidelines. Through the cytospin, BT549 cells were spotted on glass microscope slides and dried before being fixed with 4 % paraformaldehyde, and incubated with two primary antibodies raised in different species specific for the target proteins (Table 4.2). Glasses were incubated with primary antibodies diluted in PBS 3 % BSA in a humidity chamber for 2 h at RT. Then cells were stained with the PLA probes diluted 1:5 in PBS 1 % BSA in a humidified chamber for 1 h at 37 °C. Since a short DNA strand is attached to each PLA probe, when protein targets are in close proximity, an enzymatic ligation and a subsequent rolling circle amplification (at 37 °C for 100 min) occur after the addition of other circle-forming DNA oligonucleotides. At the end of the amplification step, glasses are washed
with the supplied buffers, dried and mounted with cover slips using Duo Link mounting solution containing DAPI. Glasses were stored at -20 °C until analysis, performed with fluorescence microscopy Nikon Eclipse E1000. Protein interactions result easy detectable as a distinct bright spot (Duo Link datasheet).

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Host organism</th>
<th>Dilution</th>
<th>Source</th>
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<tr>
<td>EGFR</td>
<td>mouse</td>
<td>1:200</td>
<td>Abcam</td>
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<tr>
<td>cIAP1</td>
<td>rabbit</td>
<td>1:150</td>
<td>Cell Signaling</td>
</tr>
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<td>RAB7</td>
<td>rabbit</td>
<td>1:150</td>
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<td>RAB11</td>
<td>rabbit</td>
<td>1:150</td>
<td>Cell Signaling</td>
</tr>
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</table>

Table 4.4 - List of antibodies used for PLA.

4.16 Real-Time PCR

4.16.1 RNA extraction

For Real-time qPCR, total RNA was extracted from cells with the miRNeasy mini columns (Qiagen), following the manufacturer's instructions. Briefly, cells were harvested and washed once with PBS before being lysed in QIAzol and stored at -80 °C. To proceed with the RNA extraction, 140 µl of chloroform were added to each sample and after centrifugation (15 min at 13000 rpm at 4 °C), the aqueous, upper phase, was recovered in a new microcentrifuge tube and mix with 1.5 volumes of 100 % ethanol. Then, samples were loaded into RNeasy Mini columns thus allowing the RNA isolation exploiting the affinity with the column membrane. For this, columns were centrifuged and washed one time with the RWT buffer and, then, further two times with the RPE buffer, before being transferred to a new clean and sterile tube and added with 30 µl of RNase-free water to elute RNA. Following RNA extraction, the obtained RNA was quantified using the spectrophotometer NanoDrop 2000C.
(Thermo Fisher Scientific). After quantification, 1 μg of RNA was subjected to electrophoresis (with 1 % agarose gel) to control the quality of the extracted RNA.

4.16.2 RNA reverse transcription

Transcriptor First Strand cDNA Synthesis Kit (Roche, Indianapolis, IN, USA) was used for the RNA reverse transcription to obtain and amplify cDNA in a single step. According to the manufacturer’s instructions, a mix containing 1 μg RNA and 60 μM random examers was prepared and denatured at 65 °C for 10 min. After denaturation, a mix consisting in reaction buffer (containing 8 mM MgCl₂), protector RNase inhibitor (20 U), dNTPs mix (1 mM each) and Transcriptor Reverse Transcriptase (10 U) was prepared and added to each sample to reach 20 μl final volume.

The PCR program used was:

25 °C  10 min
50 °C  50 min
85 °C  5 min

Based on the RNA used and the volume of the reaction, the PCR products (cDNA) had a concentration of 50 ng/μl. cDNA was then stored at -20 °C until Real-Time PCR analysis.

4.16.3 Quantitative Real-Time PCR

The quantification of target gene mRNA was performed using Taqman standard curve method, which specifically amplified sequences encoding the gene of interest. Probes employed for this assay (Applied Biosystems- Thermo Fisher Scientific) had FAM as fluorescent dye in the 5’ position and were conjugated to the quencher TAMRA at the 3’.
During the Real-Time PCR, gene targets are identified by specific probes and therefore amplified. The intensity of light released by the probe during this reaction is measured in real time and reflects the expression levels of the gene in the analysed sample. For each reaction the total volume was 20 µL of a mix containing 10 µl of TaqMan Universal PCR Mastermix (Applied Biosystems #4304437), 1 µl of probe (dual labelled), 4 µl of RNase-free water and 5µl of cDNA which is replaced with 5 µl of RNase-free water as a negative control (NTC). All standards, samples and negative controls were assayed in triplicate to ensure accurate results. PCR reaction was performed in 96-well plates (MicroAmp Fast Optical 96-Well Reaction Plate with Barcode, Applied Biosystems) covered with optical adhesive covers. The instrument used was ViiA™ 7 Real-Time PCR System (Applied Biosystems).

Real time PCR was performed with FAST running method:

*Hold stage*

Step 1: 95 °C 20 sec

*PCR stage (40 cycles)*

Step 1: 95 °C 1 sec

Step 2: 60 °C 20 sec

*Melting curve stage*

Step 1: 95 °C 15 sec

Step 2: 60 °C 60 sec

Step 3: 95 °C 15 sec (dissociation)

The threshold cycle (Ct Value) is the intensity of fluorescence considered statistically significant above the baseline values. Relative expression levels were calculated using the
comparative Ct method and calibrated relative to the reference gene. The probes listed in Table 4.5 were used for Real-Time PCR:

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<th>Assay ID</th>
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</thead>
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<tr>
<td>NIK</td>
<td>Hs.PT.58.3867615</td>
<td>IDT</td>
</tr>
<tr>
<td>LRIG1</td>
<td>Hs01006152_m1</td>
<td>Applied Biosystems</td>
</tr>
<tr>
<td>EGFR</td>
<td>Hs01076090_m1</td>
<td>Applied Biosystems</td>
</tr>
<tr>
<td>ZEB1</td>
<td>Hs.PT.58.3948500</td>
<td>IDT</td>
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<td>GAPDH</td>
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</table>

Table 4.5 - List of Taqman probes used for Real-Time PCR.

4.17 Statistical and image analysis

Graph and statistical analyses were performed using GraphPad Prism 5.02. For analysis of in vivo data two-tailed unpaired Student’s t-test was applied. Real-Time PCR data were analysed using two-tailed paired Student’s t-test with the date of the experiment as pairing factor. A p-value < 0.05 was considered statistically significant. Western blot densitometric analysis was performed with ImageQuant 5.2. GEP analysis was performed as previously described (Paragraph 4.6).
5 RESULTS

5.1 SM83-mediated cIAP1 depletion reduces the metastatic potential of TNBCs

5.1.1 In vivo activity of SM83 on tumour growth

The overall effect of IAP-depletion on cancer growth was investigated employing NOD/SCID mice subcutaneously engrafted with the TNBC cell line MDA-MB231. As shown in the experimental design (Figure 5.1A), when primary tumours became evident (13 days after the inoculum), animals were treated with intraperitoneal (ip) injections of SM83 (5 mg/Kg) until the end of the experiment (30 days). Notably, MDA-MB231 subcutaneous tumour volumes were reduced more than 50 % (Figure 5.1B) in SM83 treated mice compared to untreated mice, indicating an effective anti-tumour effect of SM83 administration.

![Figure 5.1](image)

Figure 5.1 - SM83 inhibits the primary tumour growth of human breast cancers in xenograft models. (A) Experimental design. (B) NOD/SCID mice engrafted subcutaneously with $5 \times 10^6$ MDA-MB231 cells were ip injected with SM83 (5 mg/Kg, 5 times/week for 3 weeks) or left untreated (4 mice/group) until the end of the experiment. (Significant differences in days 24, 27 and 30. $P = 0.0476$, 0.0391 and 0.0344, respectively. Unpaired two-tailed t test).
Aiming to evaluate the levels of SM83 targets, MDA-MB231 nodules were collected 6 h after the last SM83 injection and protein lysates were tested by western blot. As expected, SM83 administration triggers the self-ubiquitination and proteasomal degradation of cIAP1 and cIAP2, resulting in a strong reduction of cIAP1 and cIAP2 protein levels (Figure 5.2A). The treatment also affected XIAP levels and this effect is likely to be a consequence of cIAP1 down-regulation rather than being a direct effect of SM83 treatment. In fact, the silencing of cIAP1 was sufficient to induce XIAP down-regulation in MDA-MB231 cells (Figure 5.2B). Western blots failed to detect any sign of activated apoptosis (Figure 5.2C).
Figure 5.2 - cIAP1, cIAP2 and XIAP are reduced by SM83 treatment. (A) MDA-MB231 nodules were collected 6 h after the last injection and analysed by western blot. (B) MDA-MB231 cells were transfected in vitro with two siRNAs specific for cIAP1 and treated with 100 nM SM83 for 1 h. A reduction of cIAP1, cIAP2 and XIAP levels was assessed by western blot performed 72 h after transfection. Values show the fold levels of XIAP. (C) No sign of apoptosis (cleaved PARP) was detectable in MDA-MB231 primary tumours. Actin or Vinculin are shown as loading controls.
5.2 SM83 treatment reduces spontaneous lung metastasis in NOD/SCID mice xenografted with MDA-MB231 cells

Since it has been shown that IAP can play a role in migration and invasion (Fulda, 2014c), I hypothesised that SM83 could affect the metastatic process. In order to answer this question, NOD/SCID mice subcutaneously engrafted with MDA-MB231 cells were treated for three weeks with SM83 and then lungs were collected after further two weeks (Figure 5.3A).

To investigate the SM83 effect on tumours and metastatic sites, at the end of the experiment, primary tumours, lymph nodes and lungs were collected and either lysed for RNA extraction and biochemical analysis, or paraffin-embedded for IHC. First, a delay of primary tumour growth was assessed after SM83 injection (Figure 5.3B), in accordance to my previous observation (Figure 5.1B). The inhibitory effect of SM83 on the growth of primary tumours was confirmed also employing intravenous (iv) injections. As shown in Figure 5.3B, the administration route (ip vs iv) did not influence drug efficacy and, moreover, the primary tumours of treated mice started growing as well as the untreated ones after the interruption of the injections. Importantly, a significant reduction in the number and size of lung metastasis was found in SM83 treated mice bearing MDA-MB231 tumours, which are known to spontaneously metastasize to lungs in NOD/SCID mice (Minn et al., 2005; Figure 5.3A-B).
Figure 5.3 - Treatment with SM83 affects metastasis formation in NOD/SCID mice engrafted with the highly metastatic MDA-MB231 cell line. (A) Experimental design. (B) NOD/SCID mice were engrafted subcutaneously with MDA-MB231 cells and, after two weeks, were treated for 3 weeks with ip and iv injections of SM83 (5 mg/Kg, 5 times/week) in two independent experiments. Mice were killed 2 weeks after the last injection. Graph shows tumour volumes. (C) Number (untreated n = 7, SM83-treated mice n = 8; sum of two independent
experiments shown in Figure 5.1B and 5.3B; \( P = 0.0238 \). Unpaired two-tailed t test) and (D) size (35 metastases/group; \( P = 0.0107 \). Unpaired two-tailed t test) of spontaneous MDA-MB231 lung metastases.

Notably, the anti-metastatic effect was evident independently from the administration routes, i.e. iv and ip injections. Detection of micro- and macro-metastasis was allowed by the high expression of human Vimentin of MDA-MB231 (Figures 5.4A-C).
Figure 5.4 - SM83 treatment displays anti-metastasis activity independently from the administration route.

(A) Lungs of NOD/SCID mice bearing MDA-MB231 tumours were collected 2 weeks after the last injection with SM83, formalin-fixed and paraffin-embedded. IHC of lungs with anti-human Vimentin antibody. (B) The graph
shows the number of lung metastases (Untreated vs ip P = 0.0238, Untreated vs iv P = 0.0190, ip vs iv not significant; Unpaired two-tailed t test), whereas the table shows the metastasis-free MDA-MB231-bearing mice. (C) SM83 injections (5 mg/Kg, 5 times/week) were performed ip and iv in two independent experiments. Lungs were collected from NOD/SCID mice xenografted with MDA-MB231 cells and metastasis were detected through anti-Vimentin staining in IHC.

5.3 SM83 treatment perturbs gene expression of MDA-MB231 tumours

Having found a significant reduction of MDA-MB231 spontaneous lung metastasis in mice treated with SM83, I hypothesised that IAP targeting could perturb the expression of genes responsible for the metastatic process. To test this hypothesis, MDA-MB231 nodules were collected 6 h after the last SM83 injection and profiled for gene expression (Figure 5.5). Interestingly, 65 genes were significantly modulated by SM83 treatment and, among them, 50 genes resulted up-regulated and 15 down-regulated, as compared to tumours collected from untreated mice. IAPs are known to be apical modulators of several receptor complexes and eventually control MAPK and NF-κB signalling pathways. Accordingly, the expression of a number of genes involved in the NF-κB signalling pathway (e.g. baculoviral IAP repeat containing 3/BIRC3, matrix metallopeptidase-9/MMP9, intercellular adhesion molecule 1/ICAM1) or NF-κB targets (e.g. TRAF1, TRAF2, NFKBIA, NFKB2, RelB) were indeed altered.
Figure 5.5 - SM83 administration perturbs the gene expression of MDA-MB231 primary tumours. MDA-MB231 primary tumours were subjected to microarray analysis. Heat map shows the expression levels of the 65 genes significantly modulated by SM83 injection compared to the control (expression values are in log2 scale. Red indicates higher expression; green lower expression).

In order to select the genes whose down-regulation could be responsible for the SM83-mediated anti-metastatic effect, each down-regulated gene was individually knocked-down in MDA-MB231 cells and wound healing assay was performed (Figure 5.6A). Importantly, I found that the depletion of SNAI2 was the most effective in reducing the capability of cells to migrate (Figure 5.6B), whereas the viability was only slightly affected (Figure 5.6C).
Figure 5.6 - Loss of SNAI2 reduces MDA-MB231 cell motility. (A) The 15 genes down-regulated in subcutaneous MDA-MB231 nodules upon treatment with SM83 were individually silenced in MDA-MB231 cells. To assess cell motility, 4x10^4 cells were seeded in Ibidi chambers and cultured overnight to perform wound-healing experiments. After the removal of the insert, images were acquired every hour in a Cell-IQ instrument and analysed with the integrated software. Graphs represent the percentage of gap closure after 24 h of migration.
and show the average of at least four independent experiments. (B) The graph shows the same experiment illustrated in Figure 5.6A, but focused on the effect of SNAI2 depletion. The percentage of gap closure was calculated after 24 h of migration for MDA-MB231 cells transfected with control (NT1) or SNAI2-specific siRNAs. (n = 4, P = 0.0033, Unpaired two-tailed t test). (C) MDA-MB231 cells transfected in the same way were also seeded in 96-well plates and viability assessed by CellTiter-Glo assay, 5 days after transfection.

Since GEP analysis revealed that SM83 administration down-regulates SNAI2, its protein level was detected in primary tumours collected from mice treated with SM83 or left untreated (Figure 5.7A). Western blot analysis confirmed the strong reduction of SNAI2 protein in primary tumours treated by SM83. Of note, LRIG1, an ubiquitin-ligase which is known to be induced by SM administration and that I found up-regulated in GEP, showed an opposite regulation by SM83 compared to SNAI2 (Figures 5.7A-B).

Figure 5.7 - SM83 treatment down-regulates SNAI2 and up-regulates LRIG1 levels in vivo. (A) The levels of SNAI2, down-regulated in GEP (Figure 5.5), and the up-regulation of LRIG1, were assessed by western blot performed on MDA-MB231 nodules. (B) The graph shows the fold levels of SNAI2 and LRIG1 measured by densitometric analysis of western blot and normalized to Actin levels. LRIG1: Untreated vs SM83 P = 0.046; SNAI2: Untreated vs SM83 P = 0.0143; Unpaired two-tailed t test.
5.4  cIAP1, but neither cIAP2 nor XIAP, controls SNAI2 expression

As SM83 targets different IAPs, I studied which one was responsible for SNAI2 down-regulation. To this end, I silenced cIAP1, cIAP2 and XIAP through siRNAs and I found that cIAP1, and not cIAP2 or XIAP depletion, was sufficient to reduce SNAI2 levels both in MDA-MB231 and BT549 cell lines (Figure 5.8A). Then, to further address these results, two different siRNAs targeting cIAP1 were employed in a panel of TNBC cell lines (MDA-MB231, BT549, MDA-MB157 and SUM159) showing a widely valid regulation of SNAI2 mediated by this IAP (Figure 5.8B). Taken together these results show that cIAP1 is responsible for sustaining the expression of the pro-metastatic gene SNAI2.
Figure 5.8 - **cIAP1 is the sole SM target responsible for supporting SNAI2 expression.** (A) SNAI2 levels were detected by western blot in MDA-MB231 cells transfected with siRNAs targeting cIAP1, cIAP2 or XIAP, and in BT549 cells silenced for cIAP1 or XIAP. After 72 h, cells were treated for further 6 h with 100 nM SM83. (B) SNAI2 protein levels were detected in MDA-MB231, BT549, MDA-MB157 and SUM159 cells knocked-down for cIAP1 using two different siRNAs.

Since SM83 killed a small percentage of cancer cells when administered in monotherapy, I checked if SNAI2 down-regulation was a consequence of SM83-related toxicity. Therefore, viability test was performed treating the same panel of TNBC cell lines...
described above (Figure 5.8B) with 200 nM SM83, thus finding that MDA-MB231 and BT549 cells were sensitive to the treatment (Figure 5.9A). Nonetheless, SNAI2 down-regulation occurred even when cells were treated with half of the SM83 dose and also after pre-treatment with inhibitors of apoptosis (z-VAD) and necroptosis (Nec-1; Figures 5.9B-C). Accordingly, I concluded that SNAI2 reduction is dependent on cIAP1 targeting and is not a side effect of SM83 toxicity.

Figure 5.9 - SNAI2 down-regulation is not a side effect of SM83 toxicity. (A) Cell viability was tested by CellTiter-Glo in the same cell lines employed in Figure 5.8B. Cells were treated with 200 nM SM83 and viability was assessed 24 h later. (B) MDA-MB231 cells were treated with half of the dose of SM83 (100 nM) employed in Figure 5.9A and, pre-treated or not for 1 h with caspase inhibitor z-VAD or RIP1 inhibitor Nec-1 (left panel). The treatment with z-VAD was performed also in BT549 cells (right panel). Western blot was performed to
detect the total levels of SNAI2 and cIAP1, and the cleaved form of Caspase-3. Values show the fold levels of SNAI2 relative to untreated cells.

5.5 SM83 dependent activation of the non-canonical NF-kB pathway correlates with SNAI2 down-regulation

These findings suggest that SM83 anti-metastasis effect could occur via the down-regulation of SNAI2. However, specific effectors directly involved in the cIAP1/SNAI2 axis have not been identified yet. Notoriously, cIAPs are components of several receptor complexes as the TNF-R superfamily, and allow the subsequent activation of TNF-R downstream pathways. Therefore, I investigated the underlying mechanisms through which cIAPs, particularly cIAP1, could promote SNAI2 expression focusing on the TNF-dependent activation of NF-kB and MAPK signalling pathways.

5.5.1 TNF-R2, but not TNF-R1, is responsible for SNAI2 expression

To assess if the activity of TNF-Rs promotes SNAI2 expression, the endogenous ligand TNF and its receptors, TNF-R1 and 2, were knocked-down through specific siRNAs in MDA-MB231 and BT549 cell lines. Western blot analysis revealed that in both cell lines a strong down-regulation of SNAI2 occurred after TNF and TNF-R2 depletion, while TNF-R1 seems to play only a marginal role in SNAI2 expression (Figure 5.10).
Depletion of TNF-R2 affects SNAI2 expression. MDA-MB231 and BT549 cells were transfected with siRNAs targeting TNF, TNF-R1 and TNF-R2. Cells were harvested 72 h after transfection and western blot was performed to detect SNAI2 levels. Actin is shown as a loading control.

Consequently, to investigate the mechanisms through which cIAP1 promotes SNAI2 expression, I examined the role of the non-canonical NF-κB pathway whose activation is thought to be triggered by SM-mediated IAP depletion (Gyrd-Hansen and Meier, 2010). Therefore, MDA-MB231 cells were treated with SM83 for 6 h after being knocked-down for different NF-κB-related effectors (Figure 5.11A). In accordance to my previous findings, SNAI2 levels were reduced by the treatment, while protein stability was increased by NIK and NF-κB2 depletion upon SM83 treatment, thereby suggesting the non-canonical NF-κB pathway as a possible suppressor of SNAI2. To detail the role of cIAP1 in the regulation of the non-canonical NF-κB pathway, Real-Time PCR was performed in two different TNBC cell lines, MDA-MB231 and BT549. In both cases, cIAP1 depletion enhanced NIK mRNA, indicating that this IAP not only reduces the protein stability mediating ubiquitination of NIK (Zarnegar et al., 2008), but also inhibits its expression (Figure 5.11B), thus preventing the activation of the non-canonical NF-κB pathway.
Figure 5.11 - cIAP1 depletion induces NIK and consequently non-canonical NF-kB pathway. (A) Several mediators and regulators of the NF-kB pathway were knocked-down using specific siRNAs in MDA-MB231 cells. After 72 h, cells were treated with 100 nM SM83 for 6 h and harvested. Western blot shows that the depletion of NIK and NF-kB2 prevented the non-canonical NF-kB activation and blocked SM83-mediated SNAI2 down-regulation. Vinculin is shown as a loading control. (B) Real-Time PCR was performed in MDA-MB231 and BT549 cells transiently silenced for cIAP1. NIK mRNA was evaluated as fold expression change relative to GAPDH, in cells depleted for cIAP1 compared to controls. MDA-MB231 $P = 0.0464$; $n = 4$ siCtr vs sicIAP1; Paired two-tailed t test. BT549: $P = 0.0487$; $n = 4$ siCtr vs sicIAP1; Paired two-tailed t test.

Indeed, time-course experiments performed in MDA-MB231 cells treated with SM83 at different time-points confirmed the negative correlation between SNAI2 expression and NF-kB2 activation (Figure 5.12A). This result was also observed in TNBC primary tumours.
from The Cancer Genome Atlas (TCGA; Figure 5.12B). MDA-MB231 cells were also stimulated by TWEAK, a ligand of the TNF family, which preferentially activates the non-canonical NF-κB pathway. As shown in Figure 5.12C, the activation of the non-canonical NF-κB pathway seems to be followed by SNAI2 down-regulation, leading me to hypothesise a possible inhibition of SNAI2 mediated by this pathway.
Figure 5.12 - SNAI2 down-regulation correlates with the activation of the non-canonical NF-kB pathway. (A) Time-course experiments performed in MDA-MB231 cells showing the reduction of SNAI2 after 100 nM SM83 administration, along with the activation of the non-canonical NF-kB. Actin is shown as a loading control. (B) NF-kB2, which recapitulates the activation of non-canonical NF-kB pathway, negatively correlates with SNAI2 expression levels in TNBC patients from TCGA dataset (Pearson’s correlation: -0.24; p-value: 9.6e-03). Expression values are expressed as log2 counts per million transcripts. (C) Time-course experiment was performed in MDA-MB231 cell line treated with 200 ng/ml TWEAK, a specific ligand for the non-canonical NF-kB pathway activation.

However, the silencing of NF-kB2 (Figures 5.11A-5.13) by itself did not result in the expected increase of SNAI2, leading me to investigate the involvement of other pathways in SNAI2 regulation. Hence, I focused on the role of MAPKs to establish if cIAP1 supports SNAI2 expression mediating the regulation of these signalling pathways.
Figure 5.13 - The targeting of NF-κB2 does not increase SNAI2 levels. Western blot was performed to detect SNAI2 levels in BT549 cells transfected with siRNAs targeting TNF-R2 and NF-κB2. Actin is shown as a loading control.

5.6 Loss of cIAP1 prevents SNAI2 accumulation through the inhibition of MEK signalling pathways

To identify which down-stream pathway is responsible for cIAP1-dependent expression of SNAI2, MDA-MB231, BT549 and MDA-MB157 cell lines were pre-treated for 2 h with LY294002, Triciribine, UO126 and SB203580, which target PI3K, AKT, MEK and p38 respectively. As shown in Figure 5.14A, the MEK inhibitor UO126 strongly reduced SNAI2
expression in all cell lines, and the same effect was obtained through the silencing of ERK1 and 2 (Figure 5.14B), thus confirming that MAPK effectors ERK1/2 regulate SNAI2. The Figure 5.14C shows that SM83 treatment decreased SNAI2 levels and also affected ERK phosphorylation. However, ERK activation did not appear to be dependent on TNF stimulation at these time-points.

Figure 5.14 - SNAI2 expression is promoted by MAPK signalling pathway. (A) MDA-MB231, BT549 and MDA-MB157 cells were treated for 2 h with 10 μM inhibitor of PI3K (LY294002), AKT (Triciribine), MEK (UO126) and p38 (SB203580). SNAI2 levels were detected by western blot. (B) MDA-MB231 were transfected with siRNAs targeting cIAP1, ERK1 and ERK2 for 72 h. Western blot shows SNAI2 levels. cIAP1 and ERK1/2 are shown as transfection controls. (C) Time-course experiments were performed in MDA-MB231 cells pre-treated with 100
nM SM83 for 1 h and then stimulated with 50 ng/ml TNF, for different time-points. SNAI2 levels were detected together with pERK1/2. Actin is used as a loading control.

Moreover, when MDA-MB231 cells were stimulated with TNF 50 ng/ml for 15 min and 4 h (Figures 5.15A-B), SNAI2 levels did not change. Overall, these data suggest that, in this setting, TNF did not exert any significant effect on SNAI2 expression, which appeared to be induced by ERK activation. Therefore, I speculated that other exogenous stimuli could be involved.

**Figure 5.15 - ERK-mediated induction of SNAI2 is not dependent on TNF stimulation.** (A) MDA-MB231 cells were transfected with siRNA targeting cIAP1 and, after 72 h, were treated with 50 ng/ml TNF, for 15 min and 4 h, and harvested. Western blot was performed to evaluate SNAI2 levels. (B) The MDA-MB231 cell line was silenced for cIAP1 and, 72 h after transfection, cells were pre-treated for 2 h with 10 μM inhibitor of MEK (UO126) and, then, stimulated with 50 ng/ml TNF for 4 h. Western blot shows the effect of ERK inhibition and/or TNF stimulation on SNAI2. Actin is shown as a loading control.
5.7 Targeting of cIAP1 affects the EGFR down-stream pathways thus inhibiting SNAI2 expression

5.7.1 SNAI2 expression is induced in response to EGFR activation

According to several studies showing that SNAI2 is expressed in an EGF-dependent manner (Lee et al., 2008; Kusewitt et al., 2009), I observed a correlation between EGFR and SNAI2 expression in breast cancer patients from TCGA (Figure 5.16A). Moreover, in a gene expression dataset of breast cancer cell lines (Heiser et al., 2012), I noticed a significant up-regulation of SNAI2 and EGFR in TNBC cells compared to Luminal ones (Figure 5.16B). Based on these observations, SNAI2 levels were detected by western blotting in a panel of breast cancer cell lines and the highest expression of this pro-metastatic factor was found in TNBC cell lines: MDA-MB231, MDA-MB157, SUM159, BT549 and HCC1937 (Figure 5.16C). Finally, wound-healing assay showed that the TNBC cells MDA-MB231, BT549 and HCC1937 migrated faster than the Luminal T47D (Figure 5.16D), thus suggesting that high levels of SNAI2 promotes cell motility and tumour aggressiveness.
Figure 5.16 - SNAI2 expression correlates with EGFR levels. (A) Scatter plot showing the relationship between SNAI2 and EGFR expression in breast cancer primary tumours from TCGA dataset. A significant correlation was observed (Pearson’s correlation coefficient= 0.47, p-value= 1.1e-59). Expression values are expressed as log2 counts per million transcripts. (B) Box plots showing a higher expression of SNAI2 and EGFR in TNBC versus Luminal cell lines from publicly available gene expression data. (C) A panel of breast cancer cell lines was tested to compare the levels of SNAI2. TNBC: MDA-MB231, MDA-MB157, SUM159, BT549, HCC1937, SUM149; Luminal: T47D, MDA-MB361, BT474, SkBr3, MDA-MB453. These cell lines were classified in accordance to Neve et al. classification (Neve et al., 2006). (D) Motility of different cell lines expressing diverse levels of SNAI2 was tested in wound-healing experiments by seeding 4x10^4 cells in Ibidi chambers. Images were acquired every hour.
in a Cell-IQ instrument and analysed with the integrated software. The graph shows the percentage of gap closure after 12 h of migration and represents the average of at least four independent experiments. MDA-MB231 vs T47D \( P<0.0001 \); BT549 vs T47D \( P=0.0005 \); HCC1937 vs T47D \( P=0.0003 \); Unpaired two-tailed t test.

Then, BT549 cells were stimulated with two specific ligands of EGFR (EGF and TGFα) and, as expected, a significant increase of SNAI2 level was observed, in particular after 2-3 h (Figure 5.17A). Furthermore, SNAI2 accumulation was detected in non-malignant epithelial cell lines, MCF10A and HME, upon EGFR stimulation or in the presence of a constitutive activated mutant of EGFR (Figure 5.17B), supporting the EGFR-dependent expression of SNAI2. As shown in Figure 5.17C, TGFα exposure increased ERK phosphorylation, suggesting that ERK activation occurred in response to EGFR stimulation.
**Figure 5.17 - EGFR activation promotes SNAI2 expression.** (A) For time-course experiments, BT549 cells were serum-starved overnight and then stimulated with 20 ng/ml EGF and TGFα. SNAI2 levels are shown together with total and activated levels of EGFR. (B) In human mammary epithelial cell lines, both parental and mutated for EGFR, SNAI2 levels were assessed by western blot. After overnight serum starvation, cells were stimulated
with 20 ng/ml EGF for the indicated times. (C) Time-course experiments show SNAI2 levels upon TGFα stimulation. SNAI2 up-regulation is shown together with the total and activated amount of EGFR and, activated ERK. Vinculin is shown as a loading control.

Moreover, to further test that SNAI2 up-regulation resulted from EGFR activation, cells were treated with cetuximab, a monoclonal antibody that targets the EGFR extracellular domain thus impeding the binding with its ligand and consequently preventing EGFR activation. Given that EGFR signalling involves a plethora of effectors among which MAPKs, I focused on ERK pathway since my data supported its crucial role in SNAI2 regulation (Figures 5.14A-B). Western blot analysis revealed that EGFR inhibition mediated by cetuximab abolished SNAI2 accumulation by preventing the activation of ERK signalling pathway (Figures 5.18 A-B) both in MDA-MB231 and BT549 cell lines.

![Figure 5.18](image) **Figure 5.18 - EGFR inhibition prevents SNAI2 up-regulation.** (A) MDA-MB231 and (B) BT549 cells were pre-treated with 100 μg/ml cetuximab for 1 h, after being serum-starved for 24 h. Next, cells were stimulated with 20 ng/ml EGF for the indicated time-points and SNAI2 levels were detected by western blot, together with total and phosphorylated ERK1/2 and EGFR. Values show the fold levels of SNAI2 relative to untreated cells.
5.7.2  cIAP1 sustains the EGFR-mediated expression of SNAI2

At this step, I asked whether the loss of cIAP1 affects the EGFR-mediated regulation of ERK signalling pathway. Therefore, MDA-MB231 and BT549 cells were silenced for cIAP1 and western blot analysis was performed after EGF exposure. Notably, a strong reduction of ERK activation could be detectable also at basal conditions in both cell lines (Figures 5.19A-B), and, interestingly, the up-regulation of SNAI2 was impaired even after EGF and TGFα stimulation, in the absence of cIAP1 (Figure 5.19C). Moreover, SNAI2 induction was also abolished in normal mammary epithelial cells either wt or bearing mutated EGFR (Figure 5.19D), supporting the idea that cIAP1 enhances the EGFR-dependent expression of SNAI2 by favouring the activation of ERK signalling pathway.
Figure 5.19 - The targeting of cIAP1 hinders EGFR-mediated expression of SNAI2. (A) MDA-MB231 and (B) BT549 cells were knocked-down for cIAP1 through specific siRNAs and, 48 h after transfection, starved overnight. SNAI2 levels were detected by western blot in unstimulated cells or stimulated with 20 ng/ml EGF for the indicated time-points. Detection of ERK1/2 and cIAP1 levels confirms the transfection efficiency. (C) Western blot analysis was performed to evaluate SNAI2 expression in BT549 and (D) MCF10A - wt or bearing
mutated EGFR. Cells were transfected as in Figure 5.19A and stimulated with the indicated EGFR ligands (20 ng/ml).

5.7.3 clAP1 regulates SNAI2 at transcription level

Since cells silenced for clAP1 showed lower levels of SNAI2 compared to control cells, I investigated whether the absence of clAP1 causes a decrease of SNAI2 protein stability. To this aim, SNAI2 protein half-life was evaluated in MDA-MB231 cells treated with cycloheximide, an inhibitor of protein synthesis (Figure 5.20A). Although SNAI2 was less expressed in cells knocked-down for clAP1 (siclAP1), the protein was found to be more stable in these cells compared to the control (siCtr). Therefore, I evaluated if a transcriptional regulation of SNAI2 is mediated by clAP1. Real-Time PCR was performed in BT549 and MCF10A cells, and SNAI2 expression level was evaluated in clAP1-depleted cells compared to control (Figure 5.20B). Interestingly, the analysis revealed that clAP1-targeting by specific siRNAs or IAP inhibitors (Figure 5.20B and 5.5, respectively) both resulted in the reduction of SNAI2 levels. Indeed, clAP1 depletion not only affected SNAI2 expression at basal levels, but also impaired its up-regulation upon EGFR-stimulation. Hence, these results indicate that clAP1 promotes SNAI2 expression, resulting in a decrease of its protein levels in the absence of clAP1 (Figure 5.20C).
Figure 5.20 - cIAP1 regulates SNAI2 mRNA levels. (A) MDA-MB231 cells were treated with 100 μg/ml cycloheximide in the presence or absence of cIAP1 and harvested at the indicated time points. Western blot was performed to assess SNAI2 levels. cIAP1 was detected as a control of silencing efficacy, while Actin as a loading control. (B) BT549 and MCF10A cells were silenced for cIAP1 as described before. After serum starvation, cells were stimulated for 3 h with 20 ng/ml EGF and lysed to extract RNA. Real-Time PCR was performed to evaluate SNAI2 fold expression relative to GAPDH. BT549: Unstimulated siCtr vs sicIAP1 P = 0.0137, EGF 3 h siCtr vs sicIAP1 P = 0.0558; n = 3; Paired two-tailed t test. MCF10A: Unstimulated siCtr vs sicIAP1...
P = 0.0784, EGF 3 h siCtr vs sicIAP1 P = 0.0742; n = 4; Paired two-tailed t test. (C) SNAI2 levels were assessed by western blot together with EGFR total and activated levels, in BT549 and MCF10A cell lines. After 48 h from transfection with control siRNA (siCtr) or siRNAs specific for cIAP1 (sicIAP1), cells were serum-starved overnight and then stimulated with 20 ng/ml EGF for the indicated time-points.

Having demonstrated that cIAP1 supports the expression of this pro-metastasis gene, I speculated that the loss of this IAP might be responsible for the SM83 anti-metastasis effect (Figures 5.3-4). Herein, in vitro motility assay was performed to test the possible cell motility variation that might occur in the absence of cIAP1 in BT549 and in MCF10A cells, wt and carrying EGFR mutation (Figure 5.21).

![Graphs showing cell motility](image)

**Figure 5.21 - Depletion of cIAP1 reduces cell motility.** Wound healing assay was performed as described in Figure 5.6B. Motility assay was carried out using scramble-shRNA and cIAP1-shRNA transduced BT549, MCF10A and MCF10A bearing mutated EGFR cells. Graphs show the percentages of gap closure after 12 h of migration. shCtr vs shcIAP1 BT549: P= 0.0043, n=12, Paired two-tailed t test; shCtr vs shcIAP1 MCF10A: P= 0.0102, MCF10A EGFR: P= 0.0029, n= 4, Paired two-tailed t test.

Based on the reduction of in vitro cell motility (Figure 5.21) and in vivo metastasis formation (Figures 5.3-4) mediated by IAP-depletion, the effect of SM83 administration was assessed also by using a syngenic mouse model injecting the highly metastatic 4T1 murine
cells in BALB/c mice (Figure 5.22). In contrast with what I expected, SM83 administration does not affect the metastatic potential of this cell line. This result can be explained by the fact that 4T1 cells express low levels of SNAI2 (Ferrari-Amorotti et al., 2013) and therefore they metastasize in a SNAI2-independent manner. Consequently, SM83-mediated down-regulation of SNAI2 cannot impact on 4T1 metastatic potential.

Figure 5.22 - SM administration does not affect 4T1 metastatic potential. 4T1 murine cells were inoculated in syngeneic BALB/c mice that were treated with vehicle or SM83. Graph shows the number of metastatic colonies.

5.8 cIAP1 modulates not only EGFR signalling, but also its protein levels

5.8.1 EGFR levels are affected by cIAP1 depletion

The strong inhibition of ERK1/2 signalling pathway that occurred after targeting cIAP1, led me to investigate whether this IAP only promotes EGFR signalling or also regulates the receptor levels. Intriguingly, I noticed that cIAP1 depletion resulted in a marked reduction of EGFR levels in both cancer and normal mammary epithelial cell lines (Figures 5.20C and 5.23A-B), that further decreased upon EGFR stimulation. Indeed, immunofluorescence
analysis highlighted that the loss of cIAP1 strongly abrogated the number of EGFR foci induced by stimulation with its ligand, suggesting an impaired capability to transduce the receptor signals (Figure 5.23C). Having shown that cIAP1 depletion results in the reduction of EGFR protein levels, I investigated whether this is due to a direct effect of cIAP1 on receptor stability or if it is caused indirectly through the regulation of the EGFR inhibitors LRIG1 and/or c-CBL (Fry et al., 2009).

Figure 5.23 - cIAP1 depletion decreases EGFR levels. (A) BT549 and (B) MCF10A cells knocked-down for cIAP1 were stimulated with 20 ng/ml EGF under serum-starved (basal) conditions, and EGFR, cIAP1 and SNAI2 levels

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were analyzed by western blot. (C) BT549 cells were transfected with control or cIAP1-specific siRNAs before being overnight serum-starved and stimulated for 30 min with 20 ng/ml EGF. Cells were fixed and incubated with anti-EGFR antibody and nuclei stained with DAPI. Images were obtained with a 60x magnification.

5.8.2 cIAP1 targeting destabilizes endogenous EGFR in an LRIG1-independent manner

It has been shown that LRIG1 up-regulation represents an acquired mechanism for SM resistance (Bai et al., 2012). GEP and western blot analyses confirmed that LRIG1 up-regulation occurred at transcriptional and protein level (Figures 5.5 and 5.7A-B, respectively) in MDA-MB231 subcutaneous tumours treated with SM83. To determine whether LRIG1 increase is a result of cIAP1 depletion, I targeted this IAP through siRNAs and, then, I performed Real-Time PCR and western blot analyses to examine LRIG1 mRNA and protein level, respectively. Although LRIG1 resulted up-regulated both at transcriptional and protein level in the absence of cIAP1 (Figures 5.24A-B), surprisingly, LRIG1 knock-down did not significantly increase EGFR (Figure 5.24C). Furthermore, in sharp contrast with the strong reduction of the receptor levels (Figures 5.20C and 5.23B), the up-regulation of LRIG1 was not evident in MCF10A cells lacking cIAP1 (Figure 5.24D). The latter evidence strongly supports the idea that the observed reduction of EGFR does not stem from LRIG1 up-regulation.
Figure 5.24 - LRIG1 up-regulation mediated by cIAP1 has a limited effect on EGFR levels. (A) Real-Time PCR was performed in BT549 cells transfected with control (siCtr) or cIAP1-targeting (sicIAP1) siRNAs to assess LRIG1 expression levels. After 48 h from transfection, cells were serum-starved overnight and stimulated with 20 ng/ml EGF. Unstimulated siCtr vs sicIAP1 P = 0.0175, EGF 3 h siCtr vs sicIAP1 P = 0.0341; n = 3; Paired two-tailed t test. (B) Western blot shows SNAI2 and LRIG1 levels in BT549 cells silenced for cIAP1 and stimulated with 20 ng/ml EGF. cIAP1 was detected as control of the transfection efficiency. (C) EGFR levels were assessed in BT549 cells silenced for cIAP1 and LRIG1, stimulated with 20 ng/ml EGF after overnight serum starvation. (D) LRIG1 expression levels were evaluated by Real-Time PCR in MCF10A cells under the same conditions described in Figure 5.24A. Unstimulated siCtr vs sicIAP1 P = 0.4585, EGF 3 h siCtr vs sicIAP1 P = 0.5052; n = 4; Paired two-tailed t test.
5.8.3  cIAP1 interacts with EGFR and modulates its stability

Having demonstrated that the absence of cIAP1 reduces EGFR levels and that LRIG1 is only partially involved, I investigated other possible mechanisms through which cIAP1 could regulate EGFR stability. I firstly evaluated the interaction between EGFR and cIAP1 using two different approaches: PLA and co-IP. Through PLA endogenous EGFR–cIAP1 protein complexes could be visualized as a single bright spot in BT549 cells (Figure 5.25A). This result was further addressed performing co-IP of ectopically expressed Myc/Flag-tagged EGFR and endogenous cIAP1 (Figure 5.25B), confirming the interaction between EGFR and its negative regulator c-CBL, which increased after EGFR stimulation.

Figure 5.25 - EGFR physically interacts with cIAP1. (A) The cIAP1/EGFR interaction was tested in the BT549 cell line by PLA (red spots). DAPI staining was used to label cell nuclei (blue). Images were acquired with a 60x
magnification. (B) Co-IP was performed in BT549 cells stably expressing Myc/Flag-tagged EGFR. Cells were serum-starved overnight and then stimulated with 20 ng/ml EGF at the indicated time-points. Cells were lysed and EGFR was immunoprecipitated with anti-Flag antibody. The interaction of ectopic EGFR with cIAP1 and c-CBL was tested through western blot.

Next, to assess whether cIAP1 plays a role in the EGFR degradation, BT549 cells transfected with siCtr and sicIAP1 were pre-treated with chloroquine, in order to block lysosomal degradation. Chloroquine administration prevented the EGFR down-regulation that occurred in the absence of cIAP1, even after EGF exposure (Figure 5.26), thus suggesting that cIAP1 could affect the receptor lysosomal degradation.

![Figure 5.26](image)

**Figure 5.26 - cIAP1 affects the lysosome-dependent degradation of EGFR.** BT549 cells were first transfected with control or cIAP1-specific siRNAs. After 48 h from transfection, cells were serum-starved overnight and exposed to 100 µM lysosome inhibitor chloroquine for 1 h and subsequently stimulated with 20 ng/ml EGF for 3 h.

Considering that the main pathway of EGFR internalization is mediated by Clathrin, which drives the receptor to its degradation (Sigismund *et al.*, 2008), I also evaluated if cIAP1
affects Clathrin expression. Unexpectedly, Clathrin levels were reduced, rather than increased, by targeting of cIAP1, as shown in Figure 5.26. In accordance to this result, immunofluorescence analysis (Figure 5.27) highlighted the reduction of EGFR internalization rates caused by cIAP1 silencing. Furthermore, labelling cells with an antibody against Clathrin showed a decreased organization of the endocytic structures in cIAP1-depleted cells, whereas a clear signal resulted visible in siCtr cells before and after EGFR stimulation. Indeed, these observations prompted me to speculate that cIAP1 targeting does not trigger EGFR degradation but, instead increased its stability.
Figure 5.27 - The targeting of cIAP1 reduces Clathrin dependent EGFR endocytosis. BT549 cells treated with control or cIAP1 targeting siRNAs were serum-starved overnight and incubated with 20 ng/ml EGF for 15 or 30
min. After fixation, cells were analyzed for endocytosis and therefore incubated with an antibody against EGFR and Clathrin. Results are representative of at least three independent experiments. Images were acquired with a 60x magnification.

5.8.4 The targeting of cIAP1 increases EGFR protein stability

To substantiate the effect of cIAP1 on EGFR stability, I examined the receptor levels in BT549 and MCF10A cells ectopically expressing Myc-tagged EGFR. As shown in Figure 5.28A, cIAP1 depletion did not reduce EGFR protein levels, but rather increased them. Basing on this result, I further investigated the effect of cIAP1 depletion on EGFR stability. To this aim, BT549 and MCF10A cells were pre-treated with cycloheximide and stimulated using EGF for different time-points. Strikingly, cIAP1 depletion promoted the receptors stability upon EGF exposure, thus enhancing its half-life and supporting the idea that cIAP1 promotes EGFR degradation (Figure 5.28B).
Figure 5.28 - Ectopic EGFR fails to be degraded in the absence of cIAP1. (A) BT549 cells stably expressing Myc/Flag-tagged EGFR were grown in serum starvation for 24 h and then stimulated with 20 ng/ml EGF for 3 h. EGFR, SNAI2 and LRIG1 levels were assessed by western blot. Ectopic EGFR was detected with anti-Myc antibody. (B) Myc/Flag-tagged EGFR was stably expressed in BT549 and MCF10A cell lines. Cells were serum-starved overnight and pre-treated with 100 (BT549) or 50 (MCF10A) µg/ml cycloheximide for 30 min before being stimulated with 20 ng/ml EGF for the indicated times.
The effect of EGFR stabilization exerted by cIAP1 loss was further sustained by the observation obtained over-expressing an ubiquitin-ligase inactive form of cIAP1 which inhibited the degradation of EGFR (Figure 5.29), supporting the notion that the catalytic activity of cIAP1 triggers to receptor degradation.

![Figure 5.29 - cIAP1 ubiquitin-ligase activity triggers EGFR degradation](image)

**Figure 5.29 - cIAP1 ubiquitin-ligase activity triggers EGFR degradation.** BT549 cells transiently transfected with wt or mutated cIAP1 were employed to evaluate EGFR ectopic levels by western blot.

5.8.5  **c-CBL reduction mediated by the lack of cIAP1 enhances EGFR stability**

Since post-translational modifications including phosphorylation and ubiquitination are events that regulate EGFR endocytosis, the effect of cIAP1 on the ubiquitin ligase c-CBL was tested. In fact, I hypothesised that loss of cIAP1 could enhance the receptor stability through the reduction of c-CBL capability to trigger EGFR degradation. To this end, Myc-tagged EGFR was co-immunoprecipitated with c-CBL in BT549 cells stimulated with EGF for 30 min, in the presence or absence of cIAP1 (Figure 5.30A). These experiments showed that c-CBL co-immunoprecipitated with EGFR mainly upon EGF stimulation and the interaction between these two proteins was slightly reduced in cells silenced for cIAP1 (Figure 5.30A).
Conversely, c-CBL over-expression limited EGFR stabilization caused by cIAP1 depletion (Figure 5.30B).

**Figure 5.30 - cIAP1 depletion stabilizes EGFR reducing its c-CBL-mediated degradation.** (A) Ectopic EGFR was immunoprecipitated as described in Figure 5.24B from BT549 cells transfected with control and cIAP1-specific siRNAs. Total levels of c-CBL and the amount of c-CBL interacting with EGFR were evaluated by western blot. (B) Lentiviral vectors were employed to transduce MCF10A and BT549 to stably express Myc/Flag-tagged EGFR. Then, cells were further transduced to over-express c-CBL or GFP as a control, in cells depleted for cIAP1.
employing specific siRNAs. After overnight serum starvation, both cell lines were stimulated with 20 ng/ml EGF and analysed by western blot to evaluate the levels of ectopic EGFR employing a Myc-tagged specific antibody.

I then investigated the phosphorylation of EGFR at the Tyr-1045 residue, which is crucial for c-CBL/EGFR interaction upon EGF stimulation. I observed that, despite the fact that total levels of EGFR were reduced upon cIAP1 silencing (Figure 5.31, upper panel), the ratio of EGFR phosphorylated was increased (Figure 5.31, bottom panel). This suggests that the reduced amount of c-CBL impairs the cell capacity to degrade EGFR.

To further study the mechanisms through which cIAP1 controls EGFR levels, I also assessed the effect of cIAP1 depletion on the expression of another EGFR antagonist, named Sprouty1. Despite its role as an EGFR inhibitor, the suppression of Sprouty1 in TNBC correlates with inhibition of cell growth, invasion and metastasis (He et al., 2016). Notably, immunoblotting analysis performed in BT549 cells showed that Sprouty1 is considerably reduced in cIAP1-silenced cells compared to control cells (Figure 5.31, upper panel).
Figure 5.31 - cIAP1 knock-down enhances EGFR phosphorylation at the c-CBL binding site Tyr-1045. BT549 cells were silenced for cIAP1 through specific siRNAs and starved for 24 h before being stimulated with 20 ng/ml EGF for the indicated time-points. Sprouty 1, EGFR total levels and EGFR phosphorylation at the Tyr-1045 were assessed by western blot (upper panel). Densitometric analysis was performed on western blots and graphs (bottom panel) show the levels of EGFR phosphorylated in Tyr-1045 normalized by the total levels of the receptor.
5.8.6 **Possible role of cIAP1 in regulating the degradation and the recycling of EGFR**

After endocytosis, EGFR is guided to early endosomes where the receptor fate is decided: it is either recycled back to the cell surface or further led to late endosomes and lysosomes for degradation. To determine which endocytic route is affected by cIAP1, I studied the interaction between EGFR and RAB family proteins, with a particular focus on RAB11 and RAB7, which play a role in recycling and late endosomes, respectively (Ceresa, 2006). By PLA assay, I found that cIAP1 silencing slightly increases the interaction between EGFR and RAB11 (Figure 5.32A), while dramatically reducing the interaction of this receptor with RAB7 (Figure 5.32B). Taking into account that cIAP1 silencing reduces the total levels of EGFR (Figures 5.20C and 5.23A-B), these findings support the idea that cIAP1 depletion could promote EGFR recycling rather than its endocytosis in late vesicles. Nonetheless, further experiments are needed to clarify the role of cIAP1 in these processes.
Figure 5.32 - Loss of cIAP1 promotes EGFR recycling. (A) BT549 cells were transfected with control or cIAP1 siRNAs. After 48 h post-transfection, cells were serum-starved overnight and then stimulated with 20 ng/ml EGF. At the indicated time points, fixed cells were tested using PLA to assess interaction between EGFR and RAB11 or (B) RAB7. Images were acquired with a 60x magnification and are representative of three independent experiments. Graphs on the right panels show the number of foci counted in 50 cells.
5.8.7 Depletion of cIAP1 inhibits EGFR transcription

Despite the increased protein stability of EGFR, the overall effect of cIAP1 targeting was a strong reduction of the receptor levels. In light of these findings, I tested whether cIAP1 controls EGFR in a transcriptional manner and, therefore, the receptor expression levels were assessed through Real-Time PCR in BT549 and MCF10A cells, silenced or not for cIAP1. The analysis revealed a significant down-regulation of EGFR mRNA in the absence of cIAP1 (Figure 5.33), therefore suggesting that the observed downregulation of EGFR could stem, at least in part, from the reduced expression of EGFR gene.
Figure 5.3.3 - Loss of cIAP1 inhibits EGFR transcription. BT549 (upper panel) and MCF10A (bottom panel) cells were silenced for cIAP1 and stimulated with EGF (20 ng/ml) for 3 h after overnight serum starvation. Real-Time PCR was performed to quantify the levels of EGFR expression relative to GAPDH. BT549: Unstimulated siCtr vs sicIAP1 P = 0.0134, EGF 3 h siCtr vs sicIAP1 P = 0.0270; n = 3; Paired two-tailed t test. MCF10A: Unstimulated siCtr vs sicIAP1 P = 0.004, EGF 3 h siCtr vs sicIAP1 P = 0.0183; n = 4; Paired two-tailed t test.

To further investigate the mechanisms through which cIAP1 transcriptionally induced EGFR, I focused on NF-κB signalling pathway that is regulated by IAPs. I first found that silencing of p65/RelA resulted in a strong reduction of EGFR (Figure 5.34), suggesting that EGFR is expressed in a NF-κB-dependent manner. Therefore, as cIAP1 promotes the NF-κB canonical pathway, and EGFR expression is promoted by NF-κB, it is possible that the
reduced EGFR levels observed upon cIAP1 depletion is a consequence of an impaired activation of NF-kB.

Figure 5.34 - cIAP1 supports EGFR transcriptionally in an NF-kB dependent manner. MDA-MB231 cells were transfected with a control siRNA or siRNAs specific for NF-kB1, NF-kB2 and RelA and analysed by western blot to detect the levels of endogenous EGFR.

5.9 cIAP1 promotes the EGFR/c-MET cross-talk

To understand more deeply the role of cIAP1 in the regulation of EGFR activity, I focused my attention also on c-MET, a receptor which is known to cross-talk with EGFR (Mueller et al., 2010). Importantly, the synergism of EGFR and c-MET pathways is implicated in the development and progression of cancer favouring cell cycle progression, motility and metastasis. In order to examine whether cIAP1 also controls c-MET levels, BT549 were depleted for cIAP1 and stimulated with the c-MET specific ligand, HGF. Compared with
control cells, c-MET levels resulted stabilized by the loss of cIAP1 even after HGF stimulation (Figure 5.35A). Furthermore, in response to c-MET depletion, cIAP1 was significantly reduced (Figure 5.35B). Based on these preliminary data, I concluded that cIAP1 might represent a functional link also between EGFR and c-MET, and that cIAP1 levels are controlled by c-MET.

**Figure 5.35 - Loss of cIAP1 induces c-MET stabilization.** (A) BT549 cells were serum-starved overnight and stimulated with recombinant human HGF 20 ng/ml or with EGF 20 ng/ml (B) for the indicated time-points. Thereafter, cell lysates were collected and loaded for western blot with the indicated antibodies. Actin was used as a loading control.
5.10 Role of cIAP1 in the regulation of EMT-inducing factors

5.10.1 Targeting of cIAP1 increases, rather than inhibiting, ZEB1 expression

Having demonstrated that IAP-depletion perturbs the gene expression of SNAI2, which is an EMT-inducing factor (Phillips and Kuperwasser, 2014), I investigated if cIAP1 also controls the expression of other members involved in this process. Real-Time PCR performed both in MDA-MB231 and BT549 cells silenced for cIAP1 showed that this IAP did not support, but rather inhibited, the expression of another EMT-activator, named ZEB1 (Lehmann et al., 2016; Figure 5.36A). However, the expression of TWIST1 (Garg, 2013) was slightly reduced in the absence of cIAP1 in BT549 cells, as shown in Figure 5.36B, and it was not detectable in MDA-MB231 cells, as confirmed by western blot analysis (Figure 5.36C). Therefore, cIAP1 regulates in different manners the diverse EMT mediators: on the one hand this IAP promotes SNAI2 expression, on the other hand is responsible for ZEB1 inhibition.
Figure 5.36 - cIAP1 inhibits the expression of ZEB1 EMT-regulator. (A) Real-Time PCR was performed as described in Figure 36 to quantify the levels of ZEB1 in MDA-MB231 and BT549 cells. MDA-MB231: siCtr vs sicIAP1 P = 0.0121; BT549: P = 0.0713; n = 5; Paired two-tailed t test. (B) TWIST1 mRNA was evaluated by Real-Time PCR performed in MDA-MB231 and BT549 cells. MDA-MB231 siCtr vs sicIAP1: not detectable; BT549 siCtr vs sicIAP1: P = 0.1023; n = 5; Paired two-tailed t test. (C) Western blot analysis was performed to detect SNAI2,
ZEB1, TWIST1 and SOX2 protein levels, in BT549 cell knocked-down for cIAP1 compared to Ctr. Actin was employed as a loading control.

5.10.2 SNAI2 down-regulation mediated by targeting of cIAP1 does not affect the epithelial marker E-Cadherin

To clarify whether cIAP1 regulates the capability of breast cancer cells to undergo EMT, MDA-MB231 cells were stably transduced using three specific shRNAs targeting SNAI2. As shown in Figure 5.37, SNAI2 was efficiently knocked-down in cells transduced with sequences B and C. Although SNAI2 expression has been reported to strongly correlate with the loss of E-Cadherin, which is an important EMT mediator (Vergara et al., 2015), my results showed that cells depleted for SNAI2 failed to increase the epithelial marker E-Cadherin, but rather down-regulated it (Figure 5.37).

![Figure 5.37 - Effect of SNAI2 depletion on different EMT targets](image)

Figure 5.37 - Effect of SNAI2 depletion on different EMT targets. MDA-MB231 cells were stably silenced for SNAI2 employing lentiviral particles. Levels of E-Cadherin, Collagen 6A2 (COL6A2), SOX9 and SOX2 were...
detected in shcIAP1 cells compared to the scrambled (shCtr), by western blot. SNAI2 and Actin levels are used as a transduction and loading control, respectively.

To further validate the previous finding, I studied the effect of cIAP1-mediated SNAI2 down-regulation on the expression of E-Cadherin and Vimentin, which is a mesenchymal marker (Figures 5.38A-B). By using the MDA-MB231 and BT549 models, I found that the depletion of cIAP1 inhibits E-Cadherin mRNA (Figure 5.38A), whereas the mesenchymal marker is not significantly modified (Figure 5.38B).

![Figure 5.38](image)

Figure 5.38 - cIAP1 depletion down-regulates rather than increases E-Cadherin expression. (A) Real-Time PCR performed in MDA-MB231 and BT549 cells transfected with control or specific-cIAP1 siRNAs to assess E-
5.11 Targeting of SNAI2 affects the transcriptional regulators of cancer stem cells SOX2 and SOX9

Recently, it has been reported a correlation between SNAI2 and SOX9, which are key determinants of stem cell state (Luanpitpong et al., 2016), supporting the role of SNAI2 as a key regulator of human breast cancer stem cells. In light of these findings, I evaluated the effect of SNAI2 depletion on the expression of two regulators SOX2 and 9, which are both implied in tumour progression and malignancy, thus linking stem/progenitor signalling with oncogenesis in cancer. My data supported the mechanistic relationship between SNAI2 and SOX2/9 that was confirmed employing MDA-MB231 cells stably silenced for SNAI2 (shSNAI2), which showed a strong decrease of both factors in the absence of SNAI2 (Figure 5.37). Furthermore, SOX2 down-regulation occurred also in MDA-MB231 and BT549 cells depleted for cIAP1 (Figure 5.36C), supporting the existence of a regulatory axis consisting of cIAP1/SNAI2/SOX2.

5.12 Effect of cIAP1/SNAI2 axis on SM83-up-regulated genes

Notably, GEP analysis revealed that SM83 treatment up-regulated several genes, including some matrix-remodelling (MMP9) or collagen-encoding (COL6A1 and COL6A2) genes (Figure 5.5), thereby suggesting a possible role of IAPs in the remodelling of tumour extracellular matrix (ECM). Importantly, Real-Time PCR showed that the up-regulation of MMP9 observed in primary tumours collected from SM83-treated mice also occurred in
MDA-MB231 and BT549 cells depleted for cIAP1 (Figure 5.39A). Then, to identify if a link exists between SM83-mediated IAP depletion and the cIAP1-mediated SNAI2 down-regulation, I evaluated the effect of SNAI2 depletion on some of those genes resulted up-regulated by SM83 treatment. SNAI2 was stably depleted in MDA-MB231 cells using the most efficient shRNA targeting SNAI2 (shSNAI2 #B, Figure 5.37). Importantly, SNAI2 depletion enhances COL6A2 levels supporting the hypothesis that its up-regulation occurred as a result of SM83-mediated SNAI2 down-regulation (Figure 5.39B). Furthermore, shSNAI2 cells also showed an increase of PLEXIN-A1, another gene resulted up-regulated by SM83 administration. Collectively, these data document a possible role of the cIAP1/SNAI2 axis in the regulation of a number of genes found up-regulated in MDA-MB231 primary tumours collected from SM83-treated mice (Figure 5.5).
Figure 5.39 - SM83 administration increases the expression of several genes modulated by the cIAP1/SNAI2 axis. (A) Real-Time PCR performed in MDA-MB231 and BT549 cells silenced using Ctr or cIAP1-specific siRNAs to evaluate MMP9 expression. MDA-MB231: siCtr vs sicIAP1 P= 0.0312; BT549 siCtr vs sicIAP1: P= 0.2750; n = 3; Paired two-tailed t test. (B) Stable knock-down of SNAI2 in MDA-MB231 cells were analysed by western blot to assess the level of COL6A2 and PLEXIN-A1 protein levels.
Figure 5.40 - Schematic of the proposed mechanism for SM83 anti-metastatic activity.
6 DISCUSSION

In this thesis, I focused on the comprehension of the role of IAPs in regulating the metastatic process. Beyond their largely described anti-apoptosis activity, IAPs act in the modulation of several receptor signalling pathways as being components of different receptor complexes, such as TNF-R superfamily members (Varfolomeev et al., 2012) and pattern recognition receptors (PRRs; Vandenabeele and Bertrand, 2012). Therefore, IAP depletion not only triggers cancer cell death, but also perturbs numerous signalling pathways including MAPK and NF-κB cascades (Varfolomeev et al., 2012). This eventually results in the modification of the expression of several genes (Chesi et al., 2016; Beug et al., 2017). Since 2007, our laboratory has contributed to the development of a class of compounds designed to inhibit IAPs and named Smac mimetics (SMs). Due to their capability to target well-known inhibitors of apoptosis and increase the cytotoxic activity of traditional chemotherapies, SMs have raised great interest for cancer therapy and their employment is currently tested in clinical trials (Fulda, 2014a). Initially considered as specific XIAP inhibitors (Sun et al., 2007), SMs were later reported to target others members of IAP family, including cIAP1, cIAP2 and ML-IAP (Condon et al., 2014; Cossu et al., 2009a), thereby resulting in diverse activities and treatment outcomes. Nonetheless, the efficacy of SM administration is still limited and there is hence the need to fully understand the mechanisms of SM activity in order to define the right settings for their successful employment and to fully exploit the potential anti-cancer effect of this class of compounds.

In my PhD studies, I decided to investigate the role of IAPs in the metastatic process. To this purpose, I took advantage of a library of about 140 SMs, which has been developed
and characterized by our laboratory in collaboration with the University of Milan. All these compounds have been tested *in vitro* to evaluate their affinity for cIAP1, cIAP2 and XIAP, and to measure their cytotoxic activity in a panel of cancer cell lines. According to its *in vitro* activity and pharmacokinetic profile, I decided to perform my experiments with the compound SM83, also termed 9a in our previous publications (Lecis *et al.*, 2012; Lecis *et al.*, 2013). Notably, SM83 is a dimeric compound able to bind simultaneously with two BIR domains within the same IAP with nanomolar affinities. Indeed, *in vitro* tests have demonstrated a potent cytotoxic effect of SM83 in sensitive cancer cells and highlighted its capability to synergise with other compounds, thus enhancing their effect both *in vitro* and *in vivo* (Lecis *et al.*, 2012). By using our SM, in my PhD work, I provide evidence that the targeting of IAPs could be a possible approach to hinder the formation of metastasis (Figure 5.40).

My data show that SM83 administration reduces MDA-MB231 spontaneous lung metastasis (Minn *et al.*, 2005), in number and size. Although this result confirms that the targeting of IAPs can affect the aggressiveness of cancer cells as already shown by other groups, the underlying mechanism are still largely unknown. In fact, SM83 anti-metastasis effect may derive from many different mechanisms, such as direct killing of cancer cells, effect on tumour microenvironment (Lecis *et al.*, 2013) and the associated blood vessels (Witt *et al.*, 2015), as well as perturbation of gene expression. In my work, I mainly focused on the latter aspect. Indeed, by gene expression profiling of primary tumours collected from mice injected with SM83, I identified the genes whose expression was altered by treatment and suggested new potential insights regarding the role of IAPs in the metastatic process. Bioinformatic analysis showed that the *in vivo* administration of SM83 perturbed 65 genes in
primary MDA-MB231 nodules. Among the modified genes, 50 were up-regulated, whilst 15 down-regulated. In accordance with other studies and confirming the role of IAPs in the regulation of the NF-κB pathway, the majority of the perturbed genes are well-known targets of the NF-κB cascade (e.g. TRAF1, TRAF2, NFKBIA, NFKB2, RelB; Chesi et al., 2016). To identify the genes that could play a role in the pro-metastatic activity of IAPs, I employed an unbiased approach by silencing all the genes down-regulated by SM83 treatment in vivo and performing wound-healing migration assays. In this way, I demonstrated that SNAI2 down-regulation, which was a consequence of IAP-targeting, results in the reduction of cell motility and, partially, cell proliferation. In light of this result, I hypothesised that the decrease of SNAI2 levels found in treated tumours could contribute to SM83 anti-metastatic potential. Importantly, among the SM83 targets, I identified cIAP1 as the sole responsible of SNAI2 expression since only its depletion, and not XIAP or cIAP2 targeting, inhibits SNAI2 expression. Wound healing assay performed using different cell lines (BT549, MCF10A wt or bearing EGFR mutation) further supported the idea that cIAP1 promotes cell motility in various tumour cell lines therefore suggesting that this is not a cell-type specific phenomenon.

Furthermore, my results suggest a novel role of cIAP1 as a regulator of SNAI2 expression, at least in breast cancer cells. As this transcription factor is a well-known promoter of metastasis and the in vivo experiments showed a massive down-regulation of SNAI2 upon SM83 administration, these findings could provide a novel link between IAPs and the metastatic process. In fact, although SM83 administration also reduces the volume of primary tumors, and this could obviously affect the number of circulating cancer cells, this effect is only marginal and it is unlikely to be sufficient to explain the dramatically reduction
of number and size of lung metastasis detected at the end of my experiments. Moreover, the
down-regulation of SNAI2 could be itself a consequence of SM83 toxicity and I therefore
investigated this point. Nonetheless, several experiments support that SNAI2 is a direct effect
of cIAP1 targeting and not a toxicity side effect. In fact, not only SM83 administration, but
also cIAP1 silencing (confirmed by using more than one sequence) resulted in SNAI2
reduction. These findings were also confirmed by pre-treating cells with pan-caspase or
necroptosis inhibitors and validated in cell lines resistant to SM treatment at any dose.

Importantly, the capability of cIAP1 to induce SNAI2 expression via the modulation of
NF-kB and MAPK signalling has been examined. First, I noticed that a negative correlation
exists between SNAI2 and NF-kB2, which is a marker of activation of the non-canonical NF-KB
pathway. Indeed, the triggering of the non-canonical NF-kB pathway in cells treated with
SM83 or TWEAK, which is a specific activator of the non-canonical NF-kB pathway, is
paralleled by SNAI2 down-regulation. However, further analyses revealed that the major
effect on SNAI2 levels derives from the activation of ERK pathway (Chen et al., 2009).
Nevertheless, in our models the activation of ERK by TNF seemed to be only marginal and
therefore I tested the effect of other stimuli responsible for activation of ERK.

To this end, a panel of TNBC cell lines has been employed, as these cells are highly
aggressive and therefore represent a favourite model for dissecting the metastatic process.
Of note, SNAI2 is often highly expressed in invasive tumours, such as aggressive TNBC cells,
and it therefore very likely to play an important role in the aggressive phenotype of this
tumour subtype. Importantly, the role of the transcription factor SNAI2 in the metastatic
process has been widely described by several groups: SNAI2 was shown to promote
metastasis and to be linked to stem features, and its expression was demonstrated to be associated with poor prognosis and cancer relapse (Harney et al., 2009). Interestingly, my work suggests that cIAP1 supports SNAI2 expression in a broadly valid manner, excluding a cell line-specific effect. Based on my results, cIAP1 could support TNBC aggressiveness by allowing the expression of high levels of the metastasis-promoting gene SNAI2. IAPs have been shown to favour the pro-metastatic features of cancer cells independently of their capability to control apoptosis (Mehrotra et al., 2010) and therefore SM treatment could represent a possible approach to reduce SNAI2-mediated metastasis, through the reduction of its apical regulator cIAP1.

To check this hypothesis, I have studied the signalling cascade controlled by cIAP1 and focused on EGFR. In fact, hyper-activation and/or over-expression of EGFR characterize the TNBC subtype, which is also distinguished by high levels of SNAI2. Moreover, EGFR is known to control ERK signalling which promotes SNAI2 expression. Intriguingly, cIAP1 and cIAP2 have recently been shown to interact also with EGFRvIII, an oncogenic mutant expressed in glioblastoma multiforme (GBM; Puliyappadamba et al., 2013), but the biological function of this interaction is still largely unknown. In our settings, I found that loss of cIAP1 inhibits SNAI2 expression upon EGFR stimulation because of prevention of ERK activation. Therefore, my data provide new insights on EGFR regulation and show for the first time the mechanistic role of cIAP1 in mediating the EGFR-dependent expression of SNAI2 via activation of ERK signalling pathway.

Moreover, I provide evidence that the loss of cIAP1 hinders EGFR activity also by reducing its levels. In fact, the silencing of cIAP1 causes a reduction of EGFR transcription.
These findings further support the employment of SMs in cancer treatment, sustaining the idea that SM-mediated IAP depletion may attenuate EGFR signalling also preventing its expression. Importantly, it is well documented that only a small percentage of cancer cell lines are killed by SMs in monotherapy (Petersen et al., 2007), while the vast majority is resistant to these compounds independently of the dose employed. Nonetheless, later works have shown that also intrinsically sensitive cancer cells can acquire resistance and this depends on the up-regulation of LRIG1, which is triggered by SM treatment (Bai et al., 2012). This ubiquitin-ligase is a negative regulator of several receptors, including EGFR (Miller et al., 2008) and c-MET (Shattuck et al., 2007). Therefore, I investigated whether LRIG1 up-regulation could be responsible, together with the reduction of gene expression, for the reduction of EGFR observed in cells depleted for cIAP1. Paradoxically, EGFR protein stability is increased in lack of cIAP1. I then investigated whether this IAP promotes EGFR degradation by direct ubiquitination or by controlling the levels of other EGFR inhibitors, such as c-CBL. In fact, numerous studies have defined the role of c-CBL as a regulator of ligand-induced down-regulation of EGFR (Duan et al., 2003) and proved that c-CBL regulates the cascade of events that trigger EGFR endocytosis and degradation (Haglund and Dikic, 2012). Interestingly, c-CBL physically interacts with EGFR and binds to its phosphorylated Tyr-1045 residue. This event has been shown to be crucial for EGFR degradation. In fact, several works have shown that EGFR degradation is impaired when the Tyr-1045F mutant is ectopically expressed. My data showed that the Tyr-1045 residue is more phosphorylated and this should correlate with increased degradation of EGFR in the absence of cIAP1. As already mentioned, the reduction of cIAP1 was associated with increased stability of this receptor, rather than augmented degradation. Thus, I hypothesised that EGFR, even if correctly phosphorylated on Tyr-1045,
could be degraded less efficiently in the absence of cIAP1 due to a reduction of c-CBL levels or activity. In agreement with this notion, I found that the targeting of cIAP1 causes a slight reduction of both c-CBL total levels and of the portion recruited to the EGFR complex. This evidence could explain at least in part the enhanced stability of EGFR when IAPs are targeted. Altogether these results demonstrated that cIAP1 depletion on the one hand reduces the expression of EGFR gene and on the other increases its protein stability.

In my work, I also studied the role of Sprouty1, which is another negative regulator of EGFR. Despite its role as an EGFR inhibitor, loss of Sprouty1 has been recently reported to affect EGFR-mediated mesenchymal phenotype (He et al., 2016). In particular, high expression of Sprouty1 in TNBCs has been reported to be responsible for their aggressive phenotype and promote cell migration, invasion, and anchorage-dependent and -independent growth. Accordingly, Sprouty1 depletion impairs the induction of SNAIL and SNAI2 expression by EGF, and this effect is associated with increased EGFR degradation in MDA-MB231 cells. Intriguingly, my work reveals a massive down-regulation of Sprouty1 in BT549 TNBC cells in the absence of cIAP1. This evidence may provide a further explanation regarding the mechanisms through which cIAP1 controls SNAI2 expression and the observed increase of EGFR stability in cells depleted for this IAP.

Previous reports support the importance of c-CBL-dependent ubiquitination for Clathrin-mediated endocytosis of EGFR (Jiang and Sorkin, 2003; Stang et al., 2004), which is a mechanism for RTK signal attenuation, by allowing the removal of RTKs from the cell surface (Haglund and Dikic, 2012). Interestingly, by immunofluorescence analyses, I found that the EGFR foci detected in control cells upon stimulation with EGF are abrogated by cIAP1 knock-
down. This event is also accompanied by the reduction of Clathrin expression. Since the covalent addition of ubiquitin to the EGFR receptor represents the major signal which drives to EGFR endocytosis (Haglund et al., 2003), I evaluated the effect of cIAP1 E3-ubiquitin ligase activity. The ectopic expression of an ubiquitin ligase inactive form of cIAP1 resulted in the increase of EGFR levels, supporting the notion that cIAP1 catalytic activity promotes the degradation of EGFR. Moreover, the observation that cIAP1 regulates also the levels of Clathrin suggests that this IAP tightly regulates the internalization process of EGFR.

After endocytosis, EGFR can follow two main trails: it can be recycled back to the cell surface or routed to late endosomes for lysosomal degradation. Although the importance of ubiquitination in targeting activated EGFR for degradation is established, emerging evidence show that EGFR can evade the ubiquitin-mediated degradation through the recycling to the plasma membrane. Hence, my data support the idea that cIAP1 directs EGFR to degradation while preventing its recycling after receptor endocytosis. Accordingly, by PLA, I show that the interaction between EGFR and the recycling endosome marker RAB11 is not affected by the absence of cIAP1. Conversely, loss of this IAP impairs the interaction between EGFR and late endosome marker RAB7, meaning reduced receptor degradation, in this condition.

Altogether, my thesis work supports the idea that SMs could be useful in clinics for the treatment of tumours expressing high levels of EGFR, either in monotherapy or in combination with EGFR-targeted therapy (Nakai et al., 2016). Despite the fact that SNAI2 is a well-known regulator of metastasis and tumour aggressiveness, making it a promising target for anti-cancer treatment, SNAI2-specific therapy is still lacking. In fact, so far, only proof-of-concept molecules (Harney et al., 2009) or compounds targeting SNAI2 interactors have been
described (Ferrari-Amorotti et al., 2013). Therefore, the indirect inhibition of SNAI2 mediated by SM-induced IAP depletion could represent a novel strategy to prevent metastasis formation and resistance to therapy. Accordingly, a large amount of evidence supports the role of SNAI2 in promoting cell migration and dissemination to distal organs by reducing their stem-like properties and EMT features (Kao et al., 2014).

EMT is responsible for several cellular processes, including cell proliferation, programmed cell death and differentiation during early developmental stages and tissue morphogenesis. Furthermore, many works showed a role of EMT, and of its mediator SNAI2, in metastasization even if recent works have questioned the importance of EMT in metastasis formation (Fischer et al., 2015), also proposing a negligible role of SNAI2 in cancer cell metastatic properties (Ye et al., 2015). Although transcription factors such as SNAIL, SNAI2, ZEB1, and ZEB2 are usually considered as being direct inhibitors of E-Cadherin, this point is still under debate (Cano et al., 2000). In my experiments, I found no correlation between SNAI2 and E-Cadherin, suggesting that SNAI2 does not regulate it in all conditions. This could be due to the different cellular contexts in which the experiments were performed. Intriguingly, in our settings, the depletion of cIAP1 resulted in the up-regulation of another mediator of EMT, i.e. ZEB1 (Lehmann et al., 2016). This further support the notion that IAPs, and in particular cIAP1, play a role in EMT regulation and allows me to speculate that SNAI2 down-regulation is counteracted by the increase of ZEB1. E-Cadherin levels could therefore be determined by the balance between these two opposing effects.

SNAI2 also promotes CSC-like features contributing to the aggressive tumour phenotype (Luanpitpong et al., 2016). In my study, I have confirmed that SNAI2 regulates
SOX2 and SOX9 showing that the knock-down of SNAI2 reduces SOX2 and 9 protein levels. This is consistent with the down-regulation of both SOX proteins that I have found in cIAP1-depleted cells, thus highlighting the importance of cIAP1/SNAI2 axis in the regulation of SOX2 and 9. Since CSCs are critical for dissemination of tumour cells and metastases formation (Luanpitpong et al., 2016), the effect of cIAP1 depletion could result in an anti-metastasic effect mediated by the decrease of SNAI2 dependent effect on CSCs and not on the EMT process. However, further work is necessary to clarify the role of cIAP1/SNAI2 regulatory axis in breast CSCs in order to evaluate whether its targeting could represent a therapeutic opportunity for advanced and recurrent cancers.

Finally, I investigated whether the up-regulation of PLEXIN-A1 and COL6A2 genes in MDA-MB231 nodules revealed by GEP analysis may be linked to the SM-mediated down-regulation of SNAI2. In agreement with this hypothesis, my results show that cells stably depleted for SNAI2 increase PLEXIN-A1 and COL6A2 protein levels, indicating them as SNAI2 targets. The study of SNAI2 down-stream pathway represents another important field to be explored in order to evaluate the effect of cIAP1 targeting-mediated SNAI2 inhibition. For instance, the up-regulation of COL6A2 could influence tumour matrix remodelling. Conversely, the role of PLEXIN-A1 is poorly studied. It belongs to the PLEXIN receptor family and can form complexes with RTKs, such as ErbB2, VEGFR2 or c-MET, even if the molecular mechanisms controlling these multimeric receptor complexes are poorly understood (Rizzolio and Tamagnone, 2007). These receptor complexes are diversely expressed in different phases of tumour progression and invasive growth, thereby leading to the formation of signalling complexes eliciting differential (and potentially antagonistic)
pathways. Therefore, SMs could represent a new tool to investigate their regulation and activity in tumour cells.

### 6.1 Conclusions and future research

In my PhD study, cIAP1 has been defined as a crucial determinant of the EGFR/SNAI2 axis, and this finding has been validated in a panel of breast cancer cell lines, even bearing different mutations, and in normal epithelial cells, parental or with active EGFR. Moreover, my study supports the importance of IAPs, particularly of cIAP1, in tumour dissemination and contributes to the knowledge of the mechanisms underlying cancer cell metastatic potential.

The important roles of EGFR in several cancer cell processes suggest that its targeting may provide a strategy to reduce tumour aggressiveness. Therefore, several approaches have been proposed to target EGFR, whose efficacy is frustrated by various mechanisms of resistance both primary and acquired: EGFR mutation, aberrant activation of down-stream molecules, EMT and redundant kinase signalling pathway (Luo and Fu, 2014). In the last case, a well-documented mechanism of resistance to the treatment with EGFR inhibitors is c-MET activation (Boccaccio et al., 2014). Accordingly, several studies provided the proof of principle that the combination of EGFR and c-MET inhibitors could be beneficial for the treatment of those types of cancer characterized by EGFR hyper-activity. In this regard, I have started to evaluate the effect of IAP targeting also on c-MET activity aiming to investigate a possible role of cIAP1 in mediating EGFR and c-MET cross-talk. Of note, my preliminary results indicate a stabilization of c-MET in cells depleted for cIAP1 and this event could be a consequence of decreased EGFR levels. Further experiments should be performed to elucidate the role of cIAP1 in this complex scenario to identify the most efficient
combination treatment capable to reduce the mechanisms of resistance to EGFR-directed treatment and to evaluate the efficacy of SMs in clinics.

Moreover, recent findings show that SMs not only target cancer cells, but also affects a number of immune cell populations including natural killers, myeloid cells and macrophages (Lecis et al., 2013). In my work, the capability of SM83 to perturb the tumour microenvironment has not been investigated, but the comprehension of this phenomenon is of crucial importance. In this regard, my preliminary data support the notion that SM administration could affect tumour microenvironment through the up-regulation of genes encoding for ECM components. For instance, in MDA-MB231 nodules collected from NOD/SCID mice treated with SM83, I found an increase of MMP9 expression along with COL6A2 up-regulation. Of note, matrix degradation is not the only activity of MMP9, which also regulates the availability of pro-inflammatory cytokines, chemokines and other proteins therefore regulating several aspects of inflammation and immunity. Therefore, the investigation of how IAPs influence the cross-talk between tumour cells and tumour microenvironment, including stromal cells, ECM and immune cells, is crucial to develop new therapeutic approaches.
# 7 LIST OF ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ANK</td>
<td>Ankyrin repeats motif</td>
</tr>
<tr>
<td>APAF-1</td>
<td>Apoptotic protease activating factor 1</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>AVPI</td>
<td>Alanine-Valine-Proline-Isoleucine</td>
</tr>
<tr>
<td>BAK</td>
<td>Bcl-2 homologous antagonist/killer</td>
</tr>
<tr>
<td>BAX</td>
<td>Bcl-2-associated X protein</td>
</tr>
<tr>
<td>BCA</td>
<td>Bicinchoninic acid</td>
</tr>
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<td>BCL-2</td>
<td>B-cell lymphoma-2</td>
</tr>
<tr>
<td>BH3</td>
<td>Bcl-2 homology domain 3</td>
</tr>
<tr>
<td>BID</td>
<td>BH3 interacting-domain death agonist</td>
</tr>
<tr>
<td>BIR</td>
<td>Baculoviral IAP repeat</td>
</tr>
<tr>
<td>BMP</td>
<td>Bone morphogenetic protein</td>
</tr>
<tr>
<td>BRCA</td>
<td>Breast cancer susceptibility genes</td>
</tr>
<tr>
<td>BSA</td>
<td>Bovine serum albumin</td>
</tr>
<tr>
<td>CARD</td>
<td>Caspase recruitment domain</td>
</tr>
<tr>
<td>Caspases</td>
<td>Cysteine-dependent aspartate-directed proteases</td>
</tr>
<tr>
<td>cDNA</td>
<td>Complementary DNA</td>
</tr>
<tr>
<td>CHX</td>
<td>Cycloheximide</td>
</tr>
<tr>
<td>clAP</td>
<td>Cellular IAP</td>
</tr>
<tr>
<td>CIE</td>
<td>Clathrin-independent endocytosis</td>
</tr>
<tr>
<td>CK</td>
<td>Cytokeratin</td>
</tr>
<tr>
<td>CLQ</td>
<td>Chloroquine</td>
</tr>
<tr>
<td>CME</td>
<td>Clathrin-mediated endocytosis</td>
</tr>
<tr>
<td>CSC</td>
<td>Cancer stem-like cell</td>
</tr>
<tr>
<td>DAPI</td>
<td>4',6-Diamidino-2-phenylindole dihydrochloride</td>
</tr>
<tr>
<td>DED</td>
<td>Death effector domain</td>
</tr>
<tr>
<td>DD</td>
<td>Death domains</td>
</tr>
<tr>
<td>DISC</td>
<td>Death inducing signaling complex</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco's Modified Eagle's medium</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethylsulfoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>DNTPs</td>
<td>Deoxyribonucleotide triphosphates</td>
</tr>
<tr>
<td>DR</td>
<td>Death receptor</td>
</tr>
<tr>
<td>ECL</td>
<td>Enhanced chemiluminescence</td>
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<tr>
<td>EDTA</td>
<td>Ethylenediaminetetraacetic acid</td>
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<tr>
<td>EGF</td>
<td>Epidermal growth factor</td>
</tr>
<tr>
<td>EGFR</td>
<td>Epidermal growth factor receptor</td>
</tr>
<tr>
<td>EMA</td>
<td>European Medicines Agency</td>
</tr>
<tr>
<td>EMT</td>
<td>Epithelial to mesenchymal transition</td>
</tr>
<tr>
<td>ER</td>
<td>Estrogen Receptor</td>
</tr>
<tr>
<td>ERK</td>
<td>Extracellular signal-regulated kinase</td>
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<td>FADD</td>
<td>Fas-associated protein with death domain</td>
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<tr>
<td>FBS</td>
<td>Fetal bovine serum</td>
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<tr>
<td>FDA</td>
<td>Food and Drug Administration</td>
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<tr>
<td>FDR</td>
<td>False Discovery Rate</td>
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<tr>
<td>FGF</td>
<td>Fibroblast growth factor</td>
</tr>
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<td>GEP</td>
<td>Gene expression profiling</td>
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<tr>
<td>HER2</td>
<td>Human epidermal growth factor receptor 2</td>
</tr>
<tr>
<td>HEPES</td>
<td>N'-2-hydroxyethylpiperazine-N'-2 ethanesulphonic acid</td>
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<tr>
<td>HGF</td>
<td>Hepatocyte growth factor</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<td>--------------</td>
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<tr>
<td>HME</td>
<td>Human mammary epithelial</td>
</tr>
<tr>
<td>IAP</td>
<td>Inhibitor of apoptosis protein</td>
</tr>
<tr>
<td>IBM</td>
<td>IAP binding motif</td>
</tr>
<tr>
<td>ICAD</td>
<td>Inhibitor of caspase activated DNase</td>
</tr>
<tr>
<td>ICAM1</td>
<td>Intercellular adhesion molecule 1</td>
</tr>
<tr>
<td>IF</td>
<td>Immunofluorescence</td>
</tr>
<tr>
<td>IHC</td>
<td>Immunohistochemistry</td>
</tr>
<tr>
<td>IkB</td>
<td>Inhibitor of NF-kB</td>
</tr>
<tr>
<td>IKK</td>
<td>Inhibitor of NF-kB kinase</td>
</tr>
<tr>
<td>IL</td>
<td>Interleukin</td>
</tr>
<tr>
<td>IL-1R</td>
<td>Interleukin-1 receptor</td>
</tr>
<tr>
<td>ip</td>
<td>Intraperitoneal</td>
</tr>
<tr>
<td>iv</td>
<td>Intravenous</td>
</tr>
<tr>
<td>JNK/SAPK</td>
<td>c-Jun N-terminal kinase/stress-activated protein kinase</td>
</tr>
<tr>
<td>LPS</td>
<td>Lipopolysaccharides</td>
</tr>
<tr>
<td>LRIG1</td>
<td>Leucine-rich repeats and immunoglobulin-like domains protein 1</td>
</tr>
<tr>
<td>LTβR</td>
<td>Lymptoxin β-receptor</td>
</tr>
<tr>
<td>LZ</td>
<td>Leucin-zipper-like domain</td>
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<tr>
<td>MAPK</td>
<td>Mitogen-activated protein kinases</td>
</tr>
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<td>MCL-1</td>
<td>Induced myeloid leukemia cell differentiation protein</td>
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<td>mCRC</td>
<td>Metastatic colorectal cancer</td>
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<tr>
<td>MEK</td>
<td>MAPK/ERK kinase</td>
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<tr>
<td>ML</td>
<td>Mesenchymal-like</td>
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<td>ML-IAP</td>
<td>Melanoma-IAP</td>
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<td>MMP9</td>
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<td>MOMP</td>
<td>Mitochondrial outer membrane permeabilization</td>
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<td>MSL</td>
<td>Mesenchymal stem-like</td>
</tr>
<tr>
<td>mRNA</td>
<td>Messenger RNA</td>
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<tr>
<td>mTNF</td>
<td>Membrane TFN</td>
</tr>
<tr>
<td>NAIP</td>
<td>Neuronal-IAP</td>
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<tr>
<td>NEAA</td>
<td>Non essential amino acids</td>
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<tr>
<td>NEMO</td>
<td>NF-kB essential modulator</td>
</tr>
<tr>
<td>NF-kB</td>
<td>Nuclear factor kappa-light-chain-enhancer of activated B cells</td>
</tr>
<tr>
<td>NIK</td>
<td>NF-kB inducing kinase</td>
</tr>
<tr>
<td>NOD</td>
<td>Nucleotide-binding oligomerization</td>
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<td>NOD/SCID</td>
<td>Non-obese diabetic/severe combined immunodeficiency disease</td>
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<td>NSCLC</td>
<td>Non-small cell lung cancer</td>
</tr>
<tr>
<td>PARP</td>
<td>Poly-ADP ribose polymerase</td>
</tr>
<tr>
<td>Pen/Strep</td>
<td>Penicillin, Streptomycin</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate-buffered saline</td>
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<td>PBS-T</td>
<td>PBS-Tween</td>
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<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
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<td>PI3K</td>
<td>Phosphatidylinositol 4,5-bisphosphate 3-kinase</td>
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<tr>
<td>PLA</td>
<td>Proximity ligation assay</td>
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<tr>
<td>PRRs</td>
<td>Pattern recognition receptors</td>
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<td>PR</td>
<td>Progesterone receptor</td>
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<tr>
<td>PVDF</td>
<td>Polyvinylidene difluoride</td>
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<td>Rac1</td>
<td>Ras related C3 botulinum toxin substrate</td>
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<tr>
<td>RANK</td>
<td>Receptor activator for nuclear factor kappa B</td>
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<tr>
<td>RING</td>
<td>Really interesting new gene</td>
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<td>Rel homology domain</td>
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<td>Receptor interacting protein 1</td>
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<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
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<tr>
<td>rpm</td>
<td>Revolution per minute</td>
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<tr>
<td>RPMI-1640</td>
<td>Roswell Park Memorial Institute-1640</td>
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<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<tr>
<td>RT</td>
<td>Room temperature</td>
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<td>RTK</td>
<td>Receptor tyrosine kinases</td>
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<tr>
<td>sc</td>
<td>Subcutaneous</td>
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<tr>
<td>SDS</td>
<td>Sodium dodecyl sulfate</td>
</tr>
<tr>
<td>siRNA</td>
<td>Short interfering (siRNA)</td>
</tr>
<tr>
<td>Smac/DIABLO</td>
<td>Second mitochondria-derived activator of caspase/direct inhibitor of apoptosis-binding protein with low pI</td>
</tr>
<tr>
<td>SM</td>
<td>Smac mimic</td>
</tr>
<tr>
<td>sTNF</td>
<td>Soluble TNF</td>
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<td>TA</td>
<td>Transcription activation domain</td>
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<td>TAK</td>
<td>TNF-associated kinase</td>
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<tr>
<td>TCGA</td>
<td>The Cancer Genome Atlas</td>
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<tr>
<td>TGF</td>
<td>Transforming growth factor</td>
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<tr>
<td>TIM</td>
<td>TRAF-interacting motif</td>
</tr>
<tr>
<td>TLR</td>
<td>Toll-like receptors</td>
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<tr>
<td>TNBC</td>
<td>Triple negative breast cancer</td>
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<tr>
<td>TNF</td>
<td>Tumour necrosis factor</td>
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<tr>
<td>TNF-R</td>
<td>TNF-receptor</td>
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<td>TNF-RS</td>
<td>TNF-R superfamily</td>
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<td>TNF-R1-associated death domain</td>
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<tr>
<td>TRAF</td>
<td>TNF-receptor-associating factors</td>
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<td>TRAIL</td>
<td>TNF-related apoptosis-inducing ligand</td>
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<tr>
<td>TV</td>
<td>Tumour volume</td>
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<tr>
<td>TWEAK</td>
<td>TNF-related weak inducer of apoptosis</td>
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<td>UBA</td>
<td>Ubiquitin-associated domain</td>
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<td>XIAP</td>
<td>X-linked IAP</td>
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<tr>
<td>wt</td>
<td>Wild-type</td>
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<td>ZEB</td>
<td>Zinc-finger E box-binding homeobox</td>
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11 PUBLICATIONS


12 APPENDIX
Lemur tyrosine kinase 2 (LMTK2) is a determinant of cell sensitivity to apoptosis by regulating the levels of the BCL2 family members

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Using a high-throughput approach, we identified lemur tyrosine kinase 2 (LMTK2) as a novel determinant of cell sensitivity to TRAIL. LMTK2 is a poorly characterized serine/threonine kinase believed to play a role in endosomal membrane trafficking and neuronal physiology, and recently found to be mutated in diverse tumor types. We show that LMTK2 silencing sensitizes immortalized epithelial cells and cancer cells to TRAIL, and this phenomenon is accompanied by changes in the expression of BCL2 family members. In epithelial cells, LMTK2 targeting causes the down-regulation of the BCL2 and BCL-xL anti-apoptotic proteins and the reciprocal up-regulation of the pro-apoptotic protein BIM, while, in cancer cells, LMTK2 knock-down reduces BCL2 without increasing BIM levels. We provide evidence that both BIM and BCL2 proteins are regulated by LMTK2 in a GSK3β- and PP1A-dependent manner and that their perturbation, together with BCL-xL reduction, determines an increased sensitivity not only to TRAIL, but also to other compounds. Overall, our findings suggest a broad function of LMTK2 in the regulation of the apoptotic pathway and highlight LMTK2 as a novel candidate target to increase the cytotoxic activity of chemotherapeutic compounds.

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Introduction

Among the several approaches and molecules that have been proposed in anti-cancer treatment, tumor necrosis factor-related apoptosis inducing ligand (TRAIL) has raised great expectations owing to its in vitro and in vivo cytotoxic activity towards cancer cells [1,2], while sparing normal tissues [3]. Unfortunately, later studies demonstrated that cancer cells develop resistance to TRAIL-induced apoptosis through several mechanisms, thus limiting its therapeutic efficacy [2]. Indeed, high levels of the caspase-8 and caspase-3/-7/-9 inhibitors FLIP and XIAP, hyper-activation of many pro-survival pathways (e.g. NF-kB) and aberrant expression of the BCL2 family members BCL2, BCL-xL, and MCL-1, prevent TRAIL-mediated apoptosis [2]. In pathological settings, TRAIL administration could even paradoxically sustain cancer cell aggressiveness and promote invasion and metastasis formation [4,5]. Hence, cancer cells need to be primed to death with combination therapy and, to this end, several approaches have been pursued [6–8].

Lemur Tyrosine Kinase 2 (LMTK2) is a poorly characterized serine/threonine kinase initially identified as an interactor of PP1A [9], one of the four PP1 catalytic isoforms [10]. Little is known about the physiological function of LMTK2, except for its role in the maturation of germ cells [11] and in endosomal membrane trafficking through the interaction with myosin VI [12,13]. LMTK2 is a target of CDK5 [14] and has been shown to control Smad2 signaling by regulating PP1A and GSK3β [15]. The latter kinase is constitutively active in unstimulated cells [16], but can undergo a rapid and reversible repression in response to extracellular signals. The regulation of GSK3β is mainly posttranslational and, in particular, dependent on inhibitory phosphorylation. Among the
phosphorylated residues identified, Ser9 is a known target of AKT [17] while Thr43 of ERK [18].

Due to the interaction of LMTK2 with two proteins involved in numerous processes within the nervous system, i.e. PTPA and CDK5, it is conceivable that LMTK2 may play a role in neurodegeneration [14]. Nonetheless, several LMTK2 mutations have recently been identified also in cancer, and in particular in pulmonary sarcomatoid carcinomas [19], lung adenocarcinomas [20] and prostate cancer [21,22]. Of note, in vitro models of prostate cancer, LMTK2 was shown to regulate the androgen receptor (AR) [23], prostate specific antigen (PSA) and vascular endothelial growth factor (VEGF) secretion [24], and transforming growth factor beta (TGFβ) signaling [15], and could therefore contribute to cancer susceptibility and progression [21].

In this study, we performed a siRNA-based high-throughput screening, targeting 714 kinases and 598 ubiquitin-related proteins, with the ultimate goal of identifying new modulators of TRAIL sensitivity. We found that LMTK2 is a determinant of TRAIL sensitivity and described for the first time its role in the regulation of the BCL-2 family proteins and the underlying mechanisms. Finally, we investigated the effect of LMTK2 silencing in a panel of premalignant and cancer cell lines, confirming a broad effect of LMTK2 in the regulation of the apoptotic pathway triggered not only by TRAIL, but also by other cytotoxic compounds.

**Materials and methods**

**Cell culture and chemicals**

Human mammary epithelial HME and MCF10A cell lines parental or bearing the EGFRele746A750 mutation (HME EGFR and MCF10A EGFR, respectively), and colorectal HCT116 and DLD-1 cell lines were cultured as already described [25]. MDA-MB-231 and BT549 cells were cultured in RPMI 1640 medium (Lonza Group, Basel, CH) supplemented with 10% fetal bovine serum (FBS; EuroClone, Milan, IT) at 37°C and 5% CO2 in fully humidified atmosphere. Cells were treated with the NF-κB inhibitor (BAY117082; Selleck Chemicals, Munich, D); the AKT, MEK and GSK3β inhibitors (Triciribine, U0126 and AR-A014418; Enzo Life Sciences, Plymouth Meeting, PA, USA), BTAB73 (Calbiochem, Merck KGaA, Darmstadt, D) and staurosporine (Sigma– Aldrich, St Louis, MO, USA).

**High-throughput screening and viability assay**

On day 0, HME cells bearing the EGFRele746A750 mutation (named HME EGFR hereafter) were transfected with siRNAs, arrayed on 384-well plates, using a reverse transcription protocol at a final siRNA concentration of 50 nM. The siRNA pools used for the screening corresponded to the human protein kinases and ubiquitin conjugation subsets (1, 2 and 3) sub-libraries (total 1312 gene targets, 4 siRNAs per gene target; siGENOME SMARTPool technology, GE Healthcare Dharmacon, Lafayette, CO, USA). siRNA transfection was performed with Lipofectamine RNAiMAX Transfection Reagent (Thermo Fisher Scientific, Waltham, MA, USA), according to the manufacturer’s instructions. After 48 h, cells were treated with Smac mimetic (SM83) [26,27] in combination with iraconazole-irTRAIL (izTRAIL) [28] at a final concentration of 100 nM and 0.5 ng/ml respectively; a parallel set of transfected cells was left untreated. After further 48 h, cell viability was evaluated by measuring cellular ATP content, with CellTiter-Glo (Promega, Madison, WI, USA), according to manufacturer’s protocol. Two independent experiments were performed. Luminescence values were normalized to the sample median, on a per plate basis, and untreated control values were calculated for each knock-down.

Validation experiments of the 20 candidate genes identified in the screening were performed by transfecting the HME EGFR cells with the 4 individual siRNAs that composed the corresponding siRNA pools (siGENOMEx, Dharmacon). Experiments were performed in 96 well plates and viability tested using the conditions applied in the screening.

**Silencing and ectopic expression**

Silencing experiments were performed by reverse transfection using Lipofectamine RNAiMAX Transfection Reagent and siRNAs dissolved in Opti-MEM (Thermo Fisher Scientific). The final siRNA concentration was 50 nM. When a single knock-down was compared to a double knock-down, 25 nM non-targeting siRNA was added to the single targeting siRNA to reach the same final concentration in all the reactions. The individual siRNAs employed were purchased from Dharmacon (NT#1 siGENOME Non-Targeting siRNA Pool #1 D-001206-13, NT#5 siGENOME Non-Targeting siRNA # D-001210-05; LMTK2 D-003149-06 and D-003149-21; ERN1 D-004951-02 and 004951-03; CDK5 D-003239-07 and D-003239-08) and QiAGEN (siRNA/pGFP65 Hs_REL_A5; siGSK3β Hs_GSK3B_5; siPP1A Hs_PP1CA_9; siPP1R2 Hs_PP1R2_6; siPTC1 Hs_PP1CCE_5; siBIM #5 Hs_BCL2L1_5 and siBIM #15 Hs_BCL2L1_15; QAQEN, Hilden, DE). The control siRNAs (siCtrl) used in western blots and viability tests were synthesized by Eurofins Genomics (sequence 5’- CGUACGCGGAAUACUUCGATT-3’).

Caspase activity

Apoptosis induction was measured by CellEvent Caspase-3/7 Green Detection Reagent (Thermo Fisher Scientific) which was dissolved in the culturing medium 1 h before administration of izTRAIL. Caspase activity was detected in time-course experiments by fluorescence and phase contrast image acquisition using a Cell-RIQ SLF instrument (CM Technology Oy, Tampere, Finland). Green fluorescence–positive cells were identified and counted using the integrated Cell-Q Imagen software.

Western blot

To detect protein levels, cells were trypsinized, washed in 1× PBS and boiled in lysis buffer (125 mM Tris HCl pH 6.8, 5% sodium dodecyl sulfate (SDS) for 10 min. Protease and phosphatase inhibitors were added to samples, which were then sonicated and centrifuged at 11,000 rpm for 15 min at RT. Cleared supernatants were separated by SDS-PAGE on precast 4–12% Bis–Tris NuPAGE gels (Thermo Fisher Scientific) and blotted onto PVDF membranes (Merck Millipore, Darmstadt, Germany) using the XCell II blot module (Thermo Fisher Scientific). Membranes were saturated in Tris-buffered saline (TBS) with 4% BSA for 30 min and then incubated overnight with the following primary antibodies: ERN1 #3294, BAX #2772, BAD #2922, Bcl-xl #12782, PUMA #4976, BID #2002, p-p65 #3031, GSK3β #12456, pGSK3β #5558, pPA1A #2581, pPA1A #2582, Cleaved PARP #9541, Cleaved Caspase 3 #9664, phospho-Akt Ser473 #9271, Akt #2980, Phospho-MEK1/2 #9121, MEK1/2 #4694, Myc-Tag #2278 (Cell Signaling Technology, Danvers, MA, USA), ML1C st-819 (Santa Cruz Biotechnology; Dallas, TX 75220, USA), NOXA #P0180 (Merck Millipore, Darmstadt, DE), BIM #559685 (BD Biosciences; 9320 Erenbodegem, Belgium), phospho-ERK1/2 #18519, ERK1/2 #5670, LMTK2 SA84500900, Actin #A1978, Vinculin #V9131 (Sigma–Aldrich). Antibodies specific for BCL2 [29] and BCL3 [30] have been described previously. After 1 h incubation with the appropriate horse–radish peroxidase-conjugated secondary antibody (Sigma–Aldrich), proteins were detected by electrochemiluminescence (ECL) reaction (EuroClone). Band density has been calculated by ImageQuant 5.2 and normalized to Actin. An arbitrary value of 1 has been assigned to the appropriate control sample and the others have been expressed as fold values.

**Statistical analysis**

Statistical analysis and graphs were performed using GraphPad Prism 5.02. P values were calculated by paired two-tailed t-test. A value of P < 0.05 was considered statistically significant.

**Results**

**Identification of new modulators of cell sensitivity to TRAIL using a high-throughput RNAi screening approach**

To identify genes whose knock-down increases the cytotoxicity of TRAIL, we performed a siRNA-based high-throughput screening using the HME EGFR cell line. Cells were transfected with a library of human siRNAs (pools of 4 siRNAs per target gene) targeting 714 kinases and 598 ubiquitin-related proteins, and then were either left untreated or treated with SM83 in combination with izTRAIL at sub-toxic concentrations. Cell viability was determined 48 h later by measuring cellular ATP content, using a luciferase-based assay (Fig. 1a). For each target gene, sensitivity to izTRAIL/S83 treatment was assessed by comparing the viability of untreated cells with that of cells treated with izTRAIL/S83 (Fig. 1b). Genes leading to at least 3-fold reduction of viability in the presence of izTRAIL/S83 (untreated:izTRAIL/S83 ratio > 3) were selected. The siRNA pools which resulted in cell viability <80% in untreated conditions were considered toxic and excluded from further analysis. Using these criteria, we selected 20 candidate genes (Fig. 1a and c, and...
Table 1, which were further validated in deconvolution experiments using the 4 independent siRNAs that composed the siRNA pools tested initially (Supplementary Fig. S1). Genes were validated and considered true hits only if 2 or more individual siRNAs reproduced the effect observed by the siRNA pools, i.e. the ratio treated:untreated was < 0.33 (16 genes out of 20 genes were validated, green in Supplementary Fig. S1): LRSAM1, LMTK2, PRKD2, RNF183, ERN1, RAB40B, WDR24, PMVK, SYVN1, FGFR3, DTX3L, MAP2K7, BAHD1, CSNK1G1, RFN4, DTX3, CDK5, MAP4K3.

Table 1
List of the top scoring siRNA pools increasing the sensitivity of HME EGFR cells to treatment. HME EGFR cells were transfected with a library of siRNAs targeting kinases and ubiquitin-related enzymes (total 1312 gene targets; pools of 4 siRNAs per gene target) and treated with izTRAIL and SM83 combination.

<table>
<thead>
<tr>
<th>Gene ID</th>
<th>Entrez ID</th>
<th>Gene name</th>
<th>Untreated</th>
<th>SM83/TRAIL</th>
<th>Untreated vs SM83/TRAIL</th>
</tr>
</thead>
<tbody>
<tr>
<td>LRSAM1</td>
<td>90678</td>
<td>Leucine rich repeat and sterile alpha motif containing 1</td>
<td>0.914</td>
<td>0.197</td>
<td>4.640</td>
</tr>
<tr>
<td>LMTK2</td>
<td>22853</td>
<td>Lemur tyrosine kinase 2</td>
<td>1.122</td>
<td>0.265</td>
<td>4.242</td>
</tr>
<tr>
<td>PRKD2</td>
<td>25865</td>
<td>Protein kinase D2</td>
<td>0.990</td>
<td>0.224</td>
<td>4.018</td>
</tr>
<tr>
<td>RNF183</td>
<td>138065</td>
<td>Ring finger protein 183</td>
<td>0.814</td>
<td>0.206</td>
<td>3.959</td>
</tr>
<tr>
<td>LMO7</td>
<td>4008</td>
<td>LIM domain 7</td>
<td>0.812</td>
<td>0.212</td>
<td>3.830</td>
</tr>
<tr>
<td>ERN1</td>
<td>2081</td>
<td>Endoplasmic reticulum to nucleus signaling 1</td>
<td>0.866</td>
<td>0.252</td>
<td>3.752</td>
</tr>
<tr>
<td>RAB40B</td>
<td>10966</td>
<td>RAB40B, member RAS oncogene family</td>
<td>0.955</td>
<td>0.270</td>
<td>3.683</td>
</tr>
<tr>
<td>WDR24</td>
<td>84219</td>
<td>WD repeat domain 24</td>
<td>0.827</td>
<td>0.230</td>
<td>3.603</td>
</tr>
<tr>
<td>CSF1R</td>
<td>1436</td>
<td>Colony stimulating factor 1 receptor</td>
<td>0.921</td>
<td>0.256</td>
<td>3.598</td>
</tr>
<tr>
<td>PMVK</td>
<td>10654</td>
<td>Phosphomevalonate kinase</td>
<td>0.950</td>
<td>0.270</td>
<td>3.517</td>
</tr>
<tr>
<td>SYVN1</td>
<td>84447</td>
<td>Synovial apoptosis inhibitor 1, synoviolin</td>
<td>0.860</td>
<td>0.248</td>
<td>3.466</td>
</tr>
<tr>
<td>FGFR3</td>
<td>2261</td>
<td>Fibroblast growth factor receptor 3</td>
<td>0.849</td>
<td>0.250</td>
<td>3.394</td>
</tr>
<tr>
<td>DTX3</td>
<td>151636</td>
<td>Deltex 3 (Drosophila)</td>
<td>0.855</td>
<td>0.259</td>
<td>3.301</td>
</tr>
<tr>
<td>MAP2K7</td>
<td>5609</td>
<td>Mitogen-activated protein kinase kinase 7</td>
<td>0.972</td>
<td>0.297</td>
<td>3.273</td>
</tr>
<tr>
<td>BAHD1</td>
<td>22893</td>
<td>Bromo adjacent homology domain containing 1</td>
<td>0.940</td>
<td>0.288</td>
<td>3.268</td>
</tr>
<tr>
<td>CSNK1G1</td>
<td>53944</td>
<td>Casein kinase 1, gamma 1</td>
<td>0.885</td>
<td>0.272</td>
<td>3.252</td>
</tr>
<tr>
<td>RFN4</td>
<td>6047</td>
<td>Ring finger protein 4</td>
<td>0.520</td>
<td>0.287</td>
<td>3.211</td>
</tr>
<tr>
<td>DTX3</td>
<td>196403</td>
<td>Deltex homolog 3 (Drosophila)</td>
<td>1.248</td>
<td>0.402</td>
<td>3.104</td>
</tr>
<tr>
<td>CDK5</td>
<td>1020</td>
<td>Cyclin-dependent kinase 5</td>
<td>0.929</td>
<td>0.300</td>
<td>3.102</td>
</tr>
<tr>
<td>MAP4K3</td>
<td>8491</td>
<td>Mitogen-activated protein kinase kinase kinase 3</td>
<td>0.918</td>
<td>0.304</td>
<td>3.020</td>
</tr>
</tbody>
</table>
MAP2K7, CSNK1G1, DTX3, CDK5 and MAP4K3. On the contrary, LM07, CSF1R, BAHD1, and RNF4 were excluded at this stage (red in Supplementary Fig. S1).

**LMTK2 affects the expression of BIM, BCL2 and BCL-xL**

Next, we checked the effect of the silencing of the validated 16 hits on several mediators of the apoptotic pathway to infer a potential mechanism for the observed sensitization. We focused mainly on LMTK2 and CDK5, which are known to interact with each other and to play a role in some common pathways [14], and ERN1, already known to affect sensitivity to TRAIL (Fig. 2a) by causing DR5 mRNA decay. Accordingly, the silencing of ERN1 was shown to result in increased levels of TRAIL receptor [31]. As no major differences were observed upon CDK5 silencing which could explain the increased sensitivity to TRAIL, following this analysis, we selected LMTK2, a still poorly characterized kinase. In fact, the down-regulation of LMTK2 caused the reduction of the anti-apoptotic BCL2 and BCL-xL proteins, and the up-regulation of the pro-apoptotic protein BIM. The HME EGFR cells were also characterized by FACs analysis for the expression of the TRAIL receptors (R1, R2 and decoy R3 and R4), since their modulation could affect the sensitivity to TRAIL. However, only TRAIL-R2 was detectable in HME cells and its levels were not increased by LMTK2 silencing (data not shown).

As TRAIL is known to activate the NF-kB pathway [4], we checked whether this occurs also in HME EGFR cells and found that TRAIL-dependent phosphorylation of the NF-kB component p65 (Fig. 2b) was more pronounced upon LMTK2 silencing. We therefore questioned whether NF-kB could be responsible for the modulation of BIM levels, as suggested in other settings [32], but neither the treatment with an inhibitor of the NF-kB pathway (BAY117082; Fig. 2c), nor p65 knock-down (Fig. 2d), prevented the LMTK2 silencing-dependent up-regulation of BIM. Therefore, the silencing of LMTK2 results in the modulation of diverse members of the BCL2 family, causing the down-regulation of the anti-apoptotic proteins BCL2 and BCL-xL, and the accumulation of BIM in a NF-kB-independent manner.

**The sensitization effect to TRAIL upon LMTK2 targeting is independent of EGFR status and is valid also in cancer cells**

As the screening was performed by employing normal epithelial HME cells bearing oncogenic EGFR, we asked whether our observations could be validated in other cell lines and were dependent on mutated EGFR. For this purpose, LMTK2 was silenced both in the HME and MCF10A cell lines, expressing wild type or mutated EGFR, and viability was evaluated in dose–response experiments (Fig. 3a–d). In both human mammary epithelial cell lines, the depletion of LMTK2 enhanced the killing activity of TRAIL in an equivalent manner (compare Fig. 3a vs 3c, and 3b vs 3d). Moreover, the sensitization effect was independent of EGFR status as both the parental cell lines and those bearing oncogenic EGFR were killed to a similar extent upon TRAIL administration (compare Fig. 3a vs 3b, and 3c vs 3d). The finding that LMTK2 silencing enhances the sensitivity to TRAIL in immortalized mammary epithelial cell lines led us to investigate whether this effect could also be observed in fully transformed cancer cells. To this end, we selected a panel of cancer cell lines, such as triple negative breast cancers (MDA-MB231 and BT549) and colon cancers (HCT116 and DLD-1), which, together with prostate and cervical tumors, often express high levels of LMTK2. We silenced LMTK2 in all these cancer cell lines and tested their sensitivity to TRAIL. Although differing in their intrinsic sensitivity to TRAIL, all the cell lines became more sensitive to the treatment upon LMTK2 knock-down (Fig. 3e–h), suggesting that LMTK2 is a determinant of TRAIL sensitivity in multiple cell types.
cell types. Therefore, the down-regulation of LMTK2 enhances the cytotoxic effect of izTRAIL independently of EGFR mutation and is not cell-line specific.

As BIM up-regulation was observed in HME EGFR cells upon LMTK2 silencing, we checked whether this was true also in MCF10A EGFR. Interestingly, BIM was up-regulated also in these cells, while expression of BCL2 did not vary significantly (Fig. 3i). Then, we checked the effect of LMTK2 silencing in the cancer cells lines described above. In clear contrast to what was observed in HME and MCF10A EGFR cells, LMTK2 knock-down decreased the levels of BIM in MDA-MB231, HCT116 and DLD-1 cells, while in the BT549 cell line it had no effect (Fig. 3i). Nonetheless, BCL2 expression was reduced upon LMTK2 silencing in all the four cell lines tested in accordance to HME EGFR cells. Overall, LMTK2-dependent regulation of BIM is more evident in non-cancer cell lines, while other members of the BCL2 family, such as BCL2 and BCL-xL (Figs. 2a and 3i), could be regulated by LMTK2 mainly in fully transformed cancer cells.

Silencing of LMTK2 affects AKT and ERK signaling

In an attempt to understand the mechanisms by which LMTK2 regulates the above described BCL2 family members and the sensitivity to TRAIL, we investigated the activation of AKT and ERK1/2, known to be induced by TRAIL [4], in control and LMTK2-silenced cells, treated with izTRAIL alone. In fact, we observed that LMTK2 silencing enhances the cytotoxic activity of TRAIL even in the absence of SM83 and therefore we studied the function of this kinase employing TRAIL in monotherapy. LMTK2 silencing resulted in a reduction of AKT and ERK1/2 phosphorylation (Fig. 4a–b) compatible with a decrease of their activity in unstimulated conditions. To understand whether the modulation of these pathways was responsible for BIM regulation, we treated the cells with inhibitors of AKT (Triciribine) or compounds that prevent ERK activation (U0126). All of these treatments increased BIM expression upon LMTK2 silencing (Fig. 4a–b) in untreated conditions. Therefore, LMTK2 silencing reduces the activation of AKT and ERK1/2 and this correlates to increased levels of BIM. Moreover, the chemical inhibition of these kinases further contributes to the increase of BIM levels.

As LMTK2 was shown to regulate GSK3β [15] and this kinase is inhibited by AKT and ERK [17-18], we checked whether GSK3β could play a role in the regulation of BIM. Therefore, cells silenced or not for LMTK2 were treated with the GSK3β inhibitor AR-A014418 and analyzed in time-course experiments for BIM expression. In control cells, the low basal expression of BIM was weakly affected by GSK3β
inhibition with AR-A014418, but BIM expression was markedly reduced in LMTK2-silenced cells (Fig. 4c–d) which expressed higher levels of BIM (Figs. 2a and 4). Importantly, the down-regulation of BIM caused by GSK3β inhibition was paralleled by the up-regulation of BCL2 (Fig. 4c–d). Altogether, our findings support the notion that the depletion of LMTK2 results in the inhibition of the AKT and ERK pathways and promotes the accumulation of BIM and the reduction of BCL2 and BCL-xL in a GSK3β-dependent fashion.

**LMTK2 controls BIM levels through the engagement of GSK3β and PP1A**

Then, we investigated the role of PP1A, which is known to interact with LMTK2 and mediate its effect on GSK3β [15], and found that LMTK2 silencing decreases the phosphorylation of PP1A (Fig. 5a, right panel, and 5b) and this correlated with a reduction of GSK3β phosphorylation (Fig. 5a, right panel, and 5b). The latter effect of LMTK2 silencing was more pronounced after the administration of izTRAIL (Fig. 5b), which resulted in less phosphorylation of GSK3β on Ser9 (i.e. increased activation). Interestingly, the silencing of PP1A slightly augmented GSK3β phosphorylation in untreated conditions (Fig. 5a–b), but profoundly after TRAIL treatment. This suggests that PP1A promotes GSK3β activity by reducing its inhibitory phosphorylation, but also counteracts its effect on BIM. Accordingly, BIM was regulated in a positive manner by GSK3β, but negatively by LMTK2 and PP1A (Fig. 5a–b).

PP1A is not the only phosphatase that regulates BIM, since the knock-down of PP1C and PP1R2, both known to interact with LMTK2 [9,33,34], also increased BIM expression (Fig. 5c). Interestingly, PP1A silencing sensitized cells to TRAIL to the same extent as LMTK2 silencing and the double knock-down showed no additive or synergistic effect on TRAIL toxicity (Fig. 5d) suggesting that these proteins play a role in the same pathway. Altogether, our findings indicate that LMTK2 promotes the inhibitory phosphorylation of PP1A which in turn is not able to dephosphorylate GSK3β at Ser9. When LMTK2 is silenced, this results in increased activity of GSK3β,
which favors the accumulation of BIM. GSK3β and PP1A mutually regulate each other, but the former is a promoter of BIM accumulation, whereas the latter had an opposite effect.

The perturbation of the BCL2 family members caused by LMTK2 silencing contributes to the increased sensitivity to TRAIL

The observation that LMTK2 silencing causes BIM accumulation and increases the sensitivity to TRAIL, prompted us to investigate the existence of a causal relationship between these events. To this end, BIM was silenced using two different siRNAs (Fig. 6a) alone or in combination with LMTK2 knock-down, as expected, sensitized cells to TRAIL. LMTK2/BIM double knock-down partially, but significantly, protected cells from the treatment. The increased cell viability (Fig. 6a) correlated with a reduced accumulation of cleaved PARP and caspase-3 proteins (Fig. 6b), and lower activity of caspases-3 and -7, as measured by fluorometric assay (Fig. 6c). Therefore, the accumulation of BIM contributes to the increased sensitivity to TRAIL, but the regulation of other proteins such as the BCL2 family members (Fig. 2a) could also affect the sensitization.

For this reason, we also investigated the effect of BCL2 and BCL-xL, which are both reduced by siLMTK2. BCL2 and BCL-xL were ectopically expressed in HME EGF cells silenced for LMTK2 or with control siRNAs (Fig. 6d) and treated with TRAIL. In agreement with our hypothesis, the ectopic expression of both BCL2 and BCL-xL partially protected from TRAIL treatment (Fig. 6d), suggesting that their reduction upon LMTK2 targeting contributes, together with BIM up-regulation, to the increased sensitivity.

Therefore, the partial rescue effect obtained by the ectopic expression of BCL2 and BCL-xL suggests that both proteins, together with BIM, play a role in the increased sensitivity to the treatment. Nonetheless, their role could be underestimated in our experimental conditions as the transfection efficiency for over-expression was approximately 30–40% (Fig. 6e).

The sensitizing effect of LMTK2 is not limited to TRAIL

As the members of the BCL2 family are general regulators of the apoptotic pathway and are not specifically appointed to sensitivity to TRAIL, we tested whether LMTK2 silencing could affect the cytotoxic effect of other chemotherapeutic compounds. First, we treated the HME EGF cells and two cancer cell lines with the BCL2 and BCL-xL inhibitor ABT737, and found that the down-regulation of LMTK2 increases the cytotoxic effect of the treatment in at least 2 out of the 3 tested cell lines (Fig. 7a–c and Table 2). Moreover, we tested if these observations were valid also for other cytotoxic agents such as staurosporine, cisplatin and etoposide. We found that in all cases LMTK2 silencing led to increased sensitivity to treatments and reduced cell viability, with the exception of MDA-MB231 cells treated with etoposide, which were not affected by LMTK2 silencing (Table 2). Thus, LMTK2 targeting affects the cell response to several cytotoxic agents and therefore this kinase could be considered as a new determinant of sensitivity to cell death.
Discussion

In this study, we show that LMTK2 represents a novel determinant of sensitivity to TRAIL both in cancer cells and in premalignant models, and its silencing results in enhanced cytotoxic activity of different compounds. This effect is paralleled by the modulation of the BCL2 family proteins, especially BCL2 and BIM. Mechanistically, the down-regulation of LMTK2 causes the reduction of AKT and ERK1/2 activation, a decrease in the inhibitory phosphorylation of GSK3β on Ser9 and consequently an increase of its activity. Moreover, it causes the reduction of PP1A (inhibitory) phosphorylation and this event further contributes to GSK3β activation [15], allowing the phosphatase to reduce the phosphorylation state of GSK3β. Nonetheless, as PP1A is a negative regulator of BIM, its activation counterbalances the GSK3β-dependent accumulation of BIM (Reviewed in the schematic Fig. 7d). Interestingly, BIM up-regulation is observed only in premalignant models, while LMTK2 silencing does not trigger BIM accumulation in cancer cell lines, where however BCL2 is still down-regulated. This observation supports the notion that the outcome of LMTK2 targeting on BIM and BCL2 levels is the result of different effects arising from more than one pathway. This is not surprising as the BIM level, for example, strictly relies not only on its gene transcription, but also on protein stability, which is caused by both phosphorylation and de-phosphorylation events [35]. Therefore, all these mechanisms could be differently regulated in normal, premalignant and fully transformed cells. Nonetheless, the anti-apoptotic effect of LMTK2 is valid for all the cell lines tested and its silencing results in increased sensitivity to TRAIL.

Thus far, the precise physiological function of LMTK2 is unknown. Knock-out mice are viable even if infertile due to defective germ cell maturation [11]. LMTK2 is highly expressed in other tissues, such as skeletal muscle [9,11] and brain [36,37]. Notably, two LMTK2 validated interactors, CDK5 [14,37], and PP1A [9], control neuronal differentiation, synaptic plasticity and axonal transport. The interaction between LMTK2 and the catalytic and regulatory subunits of PP1 has been demonstrated by independent groups [9] and shown to result in GSK3β regulation [15]. Accordingly, we observe that PP1A de-phosphorylates and therefore activates GSK3β [15,38]. Interestingly, both PP1A and LMTK2 were shown to...
be phosphorylated by CDK5 [37,38]. Also this kinase (CDK5) was found by us among the 16 hits confirmed in validation experiments and therefore represents, together with other CDKs [6,39,40], a potential target for combination therapy with TRAIL, even if the precise mechanism by which CDK5 affects TRAIL sensitivity still needs to be established.

Besides the role of LMTK2 in the neural system, recent findings have suggested a function also in cancer cells. In fact, several mutations have been identified in patients bearing pulmonary sarcomatoid carcinomas [19], lung adenocarcinomas [20] and prostate cancer [21,22], but their biological effect is still unknown. Our findings suggest that LMTK2 could promote cancer survival by hindering the apoptotic cascade. In fact, we show that LMTK2 targeting promotes the expression of the pro-apoptotic factor BIM and the down-regulation of the anti-apoptosis BCL2 family members, BCL-xL and BCL2. As these proteins are universal regulators of the apoptotic pathway, not tightly related to TRAIL-mediated apoptosis, it is conceivable that LMTK2 is a broad regulator of cell death, rather than that selectively triggered by TRAIL or other death ligands. In agreement to this idea, the silencing of LMTK2 resulted in an increased cytotoxic effect of staurosporine, etoposide, cisplatin and especially ABT737, a well characterized inhibitor of BCL2 and BCL-xL. Nonetheless, LMTK2 does not regulate the cell sensitivity to every compound, as the treatment with gemcitabine, for example, was not affected by LMTK2 silencing (data not shown). We have found that different members of the BCL2 family are controlled by LMTK2 in a cell-type dependent manner and all of them contribute to the final sensitivity. Accordingly, the up-regulation of BIM following LMTK2 knock-down is indeed responsible for the increased sensitivity to TRAIL, even if not sufficient to trigger cell death without an apoptotic stimulus, in agreement with other works [35,41]. Moreover, BIM is not the only determinant and the

Table 2
Effect of LMTK2 silencing on the cytotoxic activity of different compounds. HME EGFR, MDA-MB231 and BT549 cells were transfected with control siRNAs and siRNAs targeting LMTK2 and treated with serial dilutions of the indicated compounds. Viability was evaluated 24 h after the treatment and IC50 calculated by Graphpad Prism on the basis of at least of 3 independent experiments.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Cisplatin IC50 μM</th>
<th>Etoposide IC50 μM</th>
<th>Staurosporine IC50 μM</th>
<th>ABT737 IC50 μM</th>
</tr>
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<tbody>
<tr>
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<td>siLMTK2</td>
<td>siCtr</td>
<td>siLMTK2</td>
<td>siCtr</td>
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<td>218</td>
<td>129</td>
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<td>586</td>
<td>186</td>
<td>228</td>
<td>240</td>
</tr>
<tr>
<td>BT549</td>
<td>154</td>
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<td>151</td>
<td>69,9</td>
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<td>165</td>
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silencing of BIM only partially rescues the viability of cells depleted for LMTK2 and treated with iTRAiL. Our evidences support the notion that also the down-regulation of BCL2 and BCL2-XL triggered by LMTK2 silencing, contributes to the overall increased sensitivity to treatment. Of note, also the rescue effect of ectopic BCL2 and BCL2-XL expression is only partial, as in the case of BIM, suggesting that the overall sensitizing effect of LMTK2 silencing is ascribable to the sum of the effects of BIM up-regulation and BCL2 and BCL2-XL down-regulation. Nonetheless, the role of BCL2 and BCL2-XL could be more substantial than shown by our results as the transfection efficiency was less than 50% and therefore, their effect could be underestimated as many cells did not express the ectopic proteins.

Notably, LMTK2 silencing sensitizes not only premalignant cells but also cancer cells to TRAIL cytotoxicity, as in the case of breast and colorectal cancers. Interestingly, in these cancer cells BIM appears not involved in the increased sensitivity as its levels do not change, but rather decline, upon LMTK2 silencing. Therefore, other mechanisms are responsible for the increased sensitivity, and BCL2 reduction could play a role in this. More work is indeed necessary to understand the different mechanisms of regulation of BIM and BCL2 in different models, but, in our work, we show that both proteins are regulated in an opposite manner by GSK3β. This observation, together with the finding that LMTK2 silencing reduces the phosphorylation, and therefore, the activation of AKT, supports the existence of a still uncharacterized survival pathway involving LMTK2, AKT and GSK3β, which affect cell viability by regulating the levels of the BCL2 family members.

In conclusion, our work supports the notion that LMTK2 could represent a new potential target for therapy and lay the basis for the development of a new class of small molecules targeting this kinase. These compounds could be exploitable both in clinic and as tool for the comprehension of the physiological role of LMTK2. Nonetheless, the observation that LMTK2 targeting sensitizes both non-cancer, even if immortalized, and cancer cells to death highlights the need of further investigation to evaluate the toxicity of a LMTK2-directed therapy in primary cells and normal tissues. Therefore, more work is necessary to understand the biological role of LMTK2 and to determine whether its inhibition could be exploited as a new approach in anti-cancer therapy.

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Conflict of interest

The authors declare no conflict of interest.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.canlet.2016.12.025.

References


Oncogenic KRAS sensitizes premalignant, but not malignant cells, to Noxa-dependent apoptosis through the activation of the MEK/ERK pathway

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ABSTRACT

KRAS is mutated in about 20-25% of all human cancers and especially in pancreatic, lung and colorectal tumors. Oncogenic KRAS stimulates several pro-survival pathways, but it also triggers the trans-activation of pro-apoptotic genes. In our work, we show that G13D mutations of KRAS activate the MAPK pathway, and ERK2, but not ERK1, up-regulates Noxa basal levels. Accordingly, premalignant epithelial cells are sensitized to various cytotoxic compounds in a Noxa-dependent manner. In contrast to these findings, colorectal cancer cell sensitivity to treatment is independent of KRAS status and Noxa levels are not up-regulated in the presence of mutated KRAS despite the fact that ERK2 still promotes Noxa expression. We therefore speculated that other survival pathways are counteracting the pro-apoptotic effect of mutated KRAS and found that the inhibition of AKT restores sensitivity to treatment, especially in presence of oncogenic KRAS. In conclusion, our work suggests that the pharmacological inhibition of the pathways triggered by mutated KRAS could also switch off its oncogene-activated pro-apoptotic stimulation. On the contrary, the combination of chemotherapy to inhibitors of specific pro-survival pathways, such as the one controlled by AKT, could enhance treatment efficacy by exploiting the pro-death stimulation derived by oncogene activation.

INTRODUCTION

KRAS is a 21 KDa protein involved in cell signal transduction belonging to the RAS subfamily, which comprises several other small GTPases endowed with GTP-hydrolyzing activity. In unstimulated conditions, GTPases are bound to GDP and display low activity, unable to trigger the down-stream signaling processes. RAS proteins require GTP to be activated and undergo rapid cycles of activation and inactivation crucial for physiological signaling [1]. Because these cascades stimulate cell growth and division, aberrant RAS signaling
can also lead to cancer. The 3 human RAS genes (HRAS, KRAS, and NRAS) are among the most prevalent drivers of human cancer, with KRAS being mutated in 20-25% of all human tumors and up to 90% in certain cancer types, e.g. pancreatic cancer [2]. In these settings, KRAS activates several down-stream effectors leading to the stimulation of the RAF/mitogen-activated protein kinase kinase/extracellular signal-regulated kinase (RAF/MEK/ERK) and phosphatidylinositol-3-kinase (PI3K) pathways.

Colorectal cancer (CRC), one of the most widespread cancer types, displays in 40% of cases KRAS activating mutations, primarily involving codon 12 or 13. Several drug combinations are currently used for CRC treatment, including oxaliplatin, 5-FU and the camptothecin (CPT) analogue irinotecan [3]. Moreover, the epidermal growth factor receptor (EGFR)-blocking antibodies cetuximab and panitumumab are approved for treatment of metastatic CRC in combination with chemotherapy and as maintenance therapy in chemorefractory tumors. Receptor tyrosine kinases such as EGFR, through the activation of the downstream GTPases, regulate MAPK and PI3K pathways. Importantly, mutations or amplification of KRAS is often associated to unresponsiveness and acquired resistance to cetuximab [4].

Even though oncogenic KRAS is often associated with poorer prognosis, its mutations have also been considered for targeted therapy taking advantage of combinations that produce a synthetic lethal effect [5, 6]. In fact, the presence of constitutively active KRAS sensitizes cancer cells to MEK and BCL-XL [7] or RAF [8] inhibition, TRAIL [9], 5-FU and oxaliplatin [10]. Nonetheless, KRAS activation is usually associated with reduced proneness to apoptosis and increased resistance to chemotherapy owing to the activation of pro-survival pathways [11-13] and resulting in the up-regulation of anti-apoptotic factors such as the members of the inhibitor of apoptosis proteins (IAP) family [14, 15].

IAPs are characterized by the presence of a conserved baculoviral IAP repeat (BIR) domain [16] important for protein-protein interactions. Despite the 8 members of the IAP family had initially been considered essentially apoptosis negative regulators, only X-linked IAP (XIAP) is known to physically interact with caspases and prevent their activity [17]. Later studies have shown that IAPs regulate cell life aspects other than apoptosis. Cellular IAP1 (cIAP1) and cIAP2, for example, modulate the signaling of pro-survival pathways, such as the ones regulated by NF-kB transcription factors and MAPKs [16]. Interestingly, IAPs are often deregulated in cancer cells and associated to unfavorable prognosis [18]. An opportunity to target IAPs, and especially cIAP1, cIAP2 and XIAP, both for therapeutic purposes and as tools in pre-clinic research is represented by second mitochondria-derived activator of caspases (SMAC) mimetic (SM) small compounds [19]. SMs were designed to mimic the activity of SMAC [20], a natural antagonist of XIAP, which, by interacting with its BIR domains, displaces caspases and promotes their activity with consequent apoptosis induction. SMs also target cIAP1 and cIAP2, causing their degradation [21, 22], modulating several pathways and overcoming cancer cell resistance to therapy [23] and especially to tumor necrosis factor-related apoptosis inducing ligand (TRAIL) [20, 24].

Here we report that SM83, a SM recently described by us [25, 26], greatly enhances the cytotoxic activity of the topoisomerase I inhibitor CPT in premalignant models in which KRAS G13D is endogenously or ectopically expressed in human epithelial cells. The increased sensitivity of oncogenic KRAS-expressing cells stems at least in part from the basal up-regulation of the pro-apoptotic protein Noxa, which is stimulated in an ERK2-dependent manner. In clear contrast to the premalignant models, a panel of CRC lines with knock-in (KI) and knock-out (KO) mutations of KRAS G13D showed that the sensitivity to treatment is independent of KRAS status. Accordingly, Noxa levels are unaffected by oncogenic KRAS expression and other pathways, such as the ones controlled by PI3K/AKT, protect cancer cells from the potentially pro-apoptotic stimulus of mutated KRAS.

RESULTS

The combination of SMs and CPT selectively kills premalignant epithelial cells bearing oncogenic KRAS

As SMs are rarely effective in monotherapy, but sensitize cancer cells to other compounds, we searched for drugs whose cytotoxicity can be efficiently enhanced by SM83 using a high-throughput cell based screening approach. HeLa cells were exposed in vitro to SM83 and izTRAIL in addition to a combined library of about 3000 FDA-approved small molecule inhibitors and cell viability assessed (see Materials and Methods). Of the 3000 small molecule inhibitors assessed, we found that the topoisomerase I inhibitor camptothecin (CPT) most profoundly enhanced the cytotoxic effect of SM83 (Table 1). In addition to the enhancing effect of CPT, we also found that different formulations of CPT such as 10-hydroxycamptothecin also enhanced the effects of SM83, further confirming that CPT can be effectively combined with SMs and TRAIL. We then asked whether this combination is more cytotoxic in a specific genetic background and treated a panel of premalignant and cancer cell lines with izTRAIL, SM83 and CPT alone or in combination (data not shown). Viability tests showed that the immortalized human epithelial (HME) cell line bearing a KI G13D mutation in the KRAS gene (D13+/+) is far more sensitive to SM83 plus CPT treatment compared to

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Figure 1: Oncogenic KRAS increases sensitivity of HME cells to DNA-damaging agents and TRAIL. (A) The parental human epithelial (HME) cell line and the isogenic cell lines with knock-in mutations in KRAS (G13D), PI3K (H1047R) and EGFR (delE746A750) were treated with varying doses of CPT alone (left panel) or in combination with 100 nM SM83 (right panel). Viabilities are shown after 24 h of treatment. (B) HME D13/+ cells were pre-incubated with DMSO, 50 μM z-VAD, 20 μM Nec-1 (left panel), 10 μg/ml Infliximab (IFX, middle panel) and 10 μg/ml Enbrel (right panel) for 1 h and subsequently treated with 100 nM SM83 and 1 μM CPT. Cell viability was determined after 24 h. (C, D) HME +/+ and HME D13/+ cells were mock treated and treated with 100 nM SM83, 1 μM CPT and with their combination for 6 h. Cells were lysed and subjected to western blot to detect the apoptosis markers cleaved PARP, caspase-3 and caspase-8 (C) and the SM targets cIAP1, cIAP2 and XIAP (D). Actin is the loading control, asterisks show the cleaved forms p17/p19 of caspase-3 and the pro-caspase p55/p57 forms of caspase-8, together with its cleaved forms p41/p43. One representative of two independent experiments is shown.
The results showed that reduced KRAS decreased the toxicity by about 50% (Figure 2A), thus confirming the involvement of KRAS in the enhanced sensitivity. Unfortunately, the lack of an antibody specific for mutant KRAS did not allow us to determine the efficiency of G13D down-regulation (Figure S2). Furthermore, the silencing also affected wild type KRAS, which might also have a protective role to the treatment. To overcome this limit, KRAS G13D was inducibly expressed in HME cells using doxycycline. Augmented levels of phosphorylated ERK1/2 (Figure 2B), a down-stream effector of KRAS, and GST-RBD pull-down experiments confirmed the increased expression of activated KRAS (Figure 2C) paralleled by an hypersensitivity to SM83/ CPT co-treatment (Figure 2D). We then repeated the experiments with another human epithelial cell line to exclude a possible cell line-specificity of our observation. MCF10A transduced with the KRAS G13D inducible vector confirmed that expression of mutant KRAS causes the phosphorylation of ERK1/2 (Figure 2E) and hypersensitivity to cell death (Figure 2F).

### Oncogenic KRAS-mediated up-regulation of Noxa sensitizes cells to SM83/CPT co-treatment

To determine the mechanisms by which oncogenic KRAS sensitizes non-tumoral cells to treatment, several cell lines expressing endogenous and ectopic KRAS G13D were analyzed by western blot for the levels of pro- and anti-apoptotic proteins of the Bcl-2 family (data not shown). In accordance to other works, we found that the presence of oncogenic KRAS considerably increases the basal levels of Noxa in untreated cells. Accordingly, HME bearing the KI G13D mutation displayed higher levels of Noxa compared to the parental cell line (Figure 3A left panel), while the basal levels of the Noxa natural antagonist Mcl-1 were not affected by oncogenic KRAS expression, but markedly dropped after CPT treatment in a SM83-independent manner. Moreover, transient induction of ectopic KRAS G13D in HME and MCF10A parental cell lines concurred to a marked increase of Noxa levels in both cell lines (Figure 3A right panels, upper and lower panel respectively). In line with these data, KRAS silencing reduced the levels of Noxa in HME KRAS G13D cells (Figure S3). Furthermore, since Mcl1 levels were reduced concurrently to Noxa up-regulation in HME cells (Figure 3A left panel), we checked whether Noxa increase was responsible for Mcl1 down-regulation. Mcl1 levels were therefore detected in KRAS G13D-induced HME cells which showed that the sole Noxa up-regulation in not sufficient to affect Mcl1 levels in untreated cells. We then analyzed by western blot HME KI D13/+ cells silenced with control or Noxa-specific siRNAs and treated with increasing concentrations of CPT (Figure 3B). Also in this case Mcl1 stability was independent of Noxa presence, suggesting that Mcl1 down-regulation stems from the treatment and not from Noxa up-regulation. To determine

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**Table 1: Best hits from the high-throughput screening.**

<table>
<thead>
<tr>
<th>Compound</th>
<th>P-value</th>
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<tbody>
<tr>
<td>10-hydroxycamptothecin</td>
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<tr>
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</tr>
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<td>Aminacrine hydrochloride</td>
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<tr>
<td>NETILMICIN SULFATE</td>
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</table>

The parental HME or to HME carrying mutations activating PI3K and EGFR (Figure 1A). Moreover, HME D13/+ cells were more sensitive to izTRAIL alone or in combination with SM83 (Figure S1 upper panels), to the topoisoasmerase II inhibitor etoposide (ETO) and to neocarzinostatin (NCS), a DNA double strand break inducer (Figure S1 lower panel), suggesting a general enhanced sensitivity to cell death more than a specific mechanism favoring CPT-mediated death. Pre-treatment with pan-caspase inhibitor z-VAD strongly supports the idea that SM83/CPT treatment kills HME D13/+ cells through an apoptotic mechanism (Figure 1B left panel). In fact, the blocking of caspases resulted in almost complete protection from the treatment, while necroptosis inhibitor Necrostatin-1 (Nec-1) showed only a negligible effect. Importantly, as TNF is known to be a pivotal player in SM-mediated cell death, HME D13/+ were also pre-treated with the TNF-specific blockers Infliximab (Figure 1B middle panel) and Enbrel (Figure 1B right panel) which both remarkably rescued cells from the treatment, confirming the involvement of TNF in the SM83/CPT cell killing. Finally, by biochemical analysis we further confirmed that SM83 strongly increases the pro-apoptotic effect of CPT, as is evident from the substantial accumulation of cleaved PARP, caspase-8 and -3 (Figure 1C). Importantly, the altered sensitivity to treatment in cells with wild type or mutated KRAS did not stem from a diverse expression of the SM known targets cIAP1, cIAP2 and XIAP (Figure 1D), which are also depleted at the same level by SM83.
whether the oncogenic KRAS-dependent accumulation of Noxa is responsible for the hypersensitivity to SM83/CPT, viability tests were performed after Noxa depletion (Figure S2). The results showed that Noxa silencing confers resistance to treatment in HME D13/+ cells (Figure 3C).

**KRAS-induced up-regulation of Noxa is mediated by ERK2**

We next investigated the mechanisms responsible for the acquired sensitivity of KRAS-mutated HME to treatment. Both parental and D13/+ HME cell lines were treated with CPT and SM83 in the presence of various inhibitors of the MAPK, AKT and PI3K pathways which can be stimulated by activated RAS. In parental cells, the administration of MEK1/2 inhibitors PD98059 and UO126, AKT inhibitor Triciribine or PI3K inhibitor LY294002 did not affect significantly the toxicity of SM83/CPT treatment (Figure 4A). In contrast, both MEK1/2 inhibitors partially protected D13/+ HME cells from SM83/CPT treatment and conferred resistance at the same degree as parental cells (Figure 4A). Having found that Noxa is a pivotal mediator of KRAS-dependent increased sensitivity to the combination (Figure 3C), we evaluated whether the MAPK pathway was responsible for the increased levels of Noxa. We found that both MEK inhibitors reduced, as expected, the levels of phosphorylated ERK1 and ERK2, and concurrently reduced the levels of Noxa (Figure 4B). Interestingly, also Mcl1 levels were slightly reduced by the MEK inhibitors, suggesting that both Noxa and Mcl1 expression is regulated by the MEK/ERK pathway. Importantly, MEK inhibition slightly reduced Noxa basal levels also in parental HME (left panel) suggesting that the MAPK

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**Figure 2: Endogenous and ectopic mutated KRAS confers sensitivity to SM83 and CPT co-treatment.** (A) HME +/- and HME D13/+ were transfected with siRNA targeting KRAS for 48 h and subsequently treated with 100 nM SM83 and 1 μM CPT. Cell viability was determined after 24 h of treatment. (B) HME pINDUCER20-Mock (Mock) and HME pINDUCER20-KRAS G13D (G13D) were incubated with doxycycline (Dox, 250 ng/ml) for 48 h, lysed and a western blot was performed. The presence of activated KRAS was determined by detection of phosphorylated ERK1/2. (C) Active GTP-RAS was purified in cells stimulated as in (B) by pull-down assay using the recombinant RBD domain of RAF1; HME D13/+ are shown as positive control for activated KRAS. (D) HME Mock and KRAS G13D were incubated with Dox (250 ng/ml) for 48 h and treated with 100 nM SM83 and 1 μM CPT. Cell viability was determined after 24 h. (E) MCF10A Mock and KRAS G13D were incubated with Dox (250 ng/ml) for the indicated time, lysed and analyzed by western blot for the detection of ERK1/2 and phosphorylated ERK1/2. Actin is shown as a loading control. (F) MCF10A Mock and KRAS G13D were incubated with Dox (250 ng/ml) for 48 h and treated with 100 nM SM83 and 0.1 μM CPT. Cell viability was determined after 24 h. One representative of two independent experiments is shown.
pathway stimulates Noxa also in physiological conditions. To understand whether MEK targets ERK1 and ERK2 both contribute to Noxa regulation, we silenced each of them in D13/+ HME cells and found that only ERK2 downregulation reduced Noxa levels, while ERK1 silencing marginally increased accumulation of Noxa (Figure 4C right panel). Again, Mcl1 was not down-regulated by Noxa accumulation, further confirming that the treatment with CPT, and not Noxa up-regulation, was responsible for Mcl1 reduction in HME cells (Figure 3A). In line with the regulation of Noxa observed in figure 4C, ERK1 silencing slightly, but significantly, enhanced the sensitivity of D13/+ HME cells to SM83/CPT treatment, while ERK2 silencing resulted in the opposite effect (Figure 4D).

**Sensitivity to SM83/CPT is independent of KRAS status in a panel of colorectal cancer cell lines**

Our findings support the notion that oncogenic KRAS can sensitize premalignant cells to SM83/CPT treatment. We then considered whether this also occurs in malignant cells, and for this reason we employed a panel of isogenic CRC cell lines where mutated KRAS is either KI (+/+ and D13/+, SW48 and Lim1215) or KO (D13/- and +/-, HCT-116 and DLD1). Surprisingly and in

![Image](image_url)
Figure 4: ERK2, but not ERK1 is responsible for KRAS-dependent Noxa-induction. (A) HME +/+ and HME D13/+ cell lines were pre-incubated with 50 μM PD98059, 25 μM UO126, 20 μM Triciribine and 20 μM LY294002 for 2 h, and then treated with 100 nM SM83 and 1 μM CPT. Cell viability was quantified after 24 h. One representative of three independent experiments is shown. (B) HME +/+ (left panel) and HME D13/+ (right panel) cell lines were treated with 50 μM PD98059, 25 nM UO126, 20 μM Triciribine and 20 μM LY294002 for 2 h, and subsequently analyzed by western blot to detect the phosphorylated forms of AKT, ERK1 and ERK2, their total levels (upper panels) or Noxa and Mcl1 (lower panels). Actin is shown as loading control. (C) HME +/+ and HME D13/+ were transiently transfected with the indicated siRNAs for 72 hours and subsequently analyzed by western blot to detect total and phosphorylated ERK1 and ERK2, and their total levels (left panel), Noxa and Mcl1 (lower panels). Actin is shown as loading control. (D) Parental HME +/+ and HME D13/+ cells were silenced for 48 h and then treated with DMSO and 100 nM SM83 plus 1 μM CPT for further 24 h. One representative of three independent experiments is shown. * P < 0.05 vs siCtr.
contrast to the premalignant settings, the sensitivity of the CRC cell lines to SM83/CPT treatment was independent of the KRAS status (Figure 5A-D). We then investigated the Noxa status, which was responsible for the increased sensitivity of normal epithelial cells bearing oncogenic KRAS, and found that its levels were unaffected by the presence of mutated KRAS (Figure 5E-H). Likewise, the levels of Noxa-antagonist Mcl1 were not repressed by the treatment with CPT and/or SM83, and further experiments confirmed an increased stability of Mcl1 in colorectal cancer compared to HME cells (Figure S3). Noxa basal levels were however higher in CRC than in HME cell lines (data not shown), suggesting that pro-apoptotic mediators can even be up-regulated in tumor cells, but there are likely other activated pathways that counterbalance the potential pro-apoptotic stimuli.

**Aberrant activation of AKT counterbalances KRAS-mediated pro-apoptotic scenario in colorectal cancer cells**

Despite the presence of mutant KRAS, our findings suggest that cancer cells are not sensitized to SM83/CPT treatment. Therefore, we hypothesized that malignant progression might have caused the deregulation of other pathways that can counterbalance the potential apoptotic effect of oncogenic KRAS. Interestingly, HCT116 and DLD1 cells bear mutated PI3K, which results in hyper-activation of the PI3K/AKT pathway, a signaling cascade known to promote cell survival. For this reason, we treated HCT116 and DLD1 cells bearing mutated and wild type KRAS with SM83/CPT after pre-treatment with inhibitors of MEK1/2, AKT and PI3K. Interestingly, and concordant to our hypothesis, AKT inhibition restored sensitivity to the treatment only in the presence of mutant KRAS (Figure 6A and 6B). Noxa levels were lowered by MAPK blocking (Figure 6C and 6D) as already observed in HME cells (Figure 4B), but were not affected by AKT inhibition, suggesting that the AKT pathway blocks the pro-death effect triggered by oncogenic KRAS in an independent fashion. Importantly, AKT inhibition sensitized to SM83/CPT treatment also CRC cell lines bearing wild type PI3K (Figure S4), further supporting the idea that AKT counteracts the pro-death stimulus deriving from oncogenic KRAS. Surprisingly, LY294002 treatment, which reduced AKT activation, did not sensitize to SM83/CPT treatment (Figure 6C and 6D). We speculate that this stems from the fact that the LY294002 inhibitor did not completely abolish AKT phosphorylation and therefore we tested the GDC-0941 PI3K inhibitor. This compound reduced the AKT activation more efficiently (Figure 6E left panel) and sensitized the HCT116 D13/- in the same way as Triciribine (Figure 6E right panel). Finally, we investigated the mechanisms by which Noxa levels are controlled in HCT116 cells and demonstrated that the findings described for HME are true also in this cancer cell line. In fact, the targeting of ERK1 by silencing enhanced the levels of Noxa, while a specific siRNA targeting ERK2 slightly reduced its expression (Figure 6F).

**DISCUSSION**

In our work, we searched for FDA-approved drugs that increase the cytotoxic activity of IAP-antagonizing compounds and death ligands. For this purpose, using a high-throughput approach, we combined SM83 [25] and izTRAIL [29] to a library of about 3000 compounds. CPT was identified several times among the best hits and validated, alone or in combination with SM83 and/or izTRAIL, in a panel of normal and cancer cell lines bearing KI and KO mutations in genes frequently mutated in cancer. The employment of isogenic cell lines with distinct point mutations is a powerful tool to comprehend the effect of oncogene activation [31] and addiction [32], and synthetic lethal interactions [27, 33] in cancer cells. With this approach, we found that the endogenous and ectopic expression of KRAS bearing the G13D mutation sensitizes normal, but not cancer cells, to CPT plus SM83 or TRAIL treatment, and to other DNA-damaging agents.

Since oncogenic KRAS stimulates the up-regulation of the pro-apoptotic protein Noxa [10, 34], we checked the occurrence of this event in premalignant cells and whether it was associated with the increased sensitivity to treatment. In both human epithelial cells HME and MCF10A, the expression of oncogenic KRAS was indeed responsible for the up-regulation of Noxa in a MEK/ERK-dependent manner and for the augmented death upon SM83/CPT treatment. Accordingly, chemical inhibition of the MEK1/2 kinases that results in prevention of ERK1/2 phosphorylation and silencing of ERK2, but not ERK1, down-regulated Noxa in HME D13/+ to levels comparable to parental HME (+/+ ) cells. Of note, ERK2 silencing slightly, but reproducibly, protected HME D13/+ cells, while ERK1 silencing even sensitized to SM83/CPT treatment and simultaneously increased Noxa basal levels especially in CRC cell lines. The observation that ERK1 and ERK2 display opposite effects in regulating Noxa levels and mediating chemotherapy responsiveness was also described in hepatocellular carcinoma cells [35]. In this case, ERK2 knockdown was responsible for increased Noxa levels after cisplatin treatment. Although we investigated Noxa basal levels in our experiments, these contrasting results strongly support that ERK1 and ERK2 mutually regulate each other [36] in a cell type-dependent manner. Surprisingly, MEK inhibitors strongly prevented treatment cytotoxicity, while siRNA targeting ERK1 and ERK2 only have a modest effect, despite siERK2 efficiently down-regulated Noxa levels. This suggests that other unknown regulatory mechanisms between the two ERK proteins eventually impact on the treatment outcome.

We then asked whether our findings are true not only
Figure 5: Oncogenic mutation of KRAS does not confer sensitivity to combined SM83/CPT nor stimulates Noxa levels. SW48 +/- and SW48 D13/+ (A), HCT116 +/- and HCT116 D13/- (B), Lim1215 +/- and Lim1215 D13/+ (C), DLD1 D13/- and DLD1 +/- (D) cell lines were treated with DMSO and the combination of SM83 and CPT at varying concentrations. Cell viability was evaluated after 24 h. One representative of three independent experiments is shown. SW48 +/- and SW48 D13/+ (E), HCT116 +/- and HCT116 D13/- (F), Lim1215 +/- and Lim1215 D13/+ (G), DLD1 D13/- and DLD1 +/- (H) cell lines were treated with DMSO and 100 nM SM83, 0.1 µM CPT either alone or in combination for 6 h. Cells were lysed and analyzed by western blotting to determine Noxa and Mcl1 levels. One representative of two independent experiments is shown. Asterisk indicates the specific band of Actin shown as loading control.
Figure 6: Aberrant AKT activation protects HCT116 and DLD1 cells from the pro-death effect of oncogenic KRAS.
(A) HCT116 and (B) DLD1 cells were pre-incubated with 50 μM PD98059, 25 μM UO126, 20 μM Triciribine and 20 μM LY294002 for 2 h, and then mock-treated or treated with 100 nM SM83 and 1 μM CPT. Cell viability was quantified after 24 h. (C) HCT116 and (D) DLD1 cells were treated for 2 h with the indicated inhibitors as in (A) and then analyzed by western blot to detect Noxa levels and total and phosphorylated AKT and ERK levels. Actin is shown as loading control. (E, left panel) HCT116 D13/- cells were treated with 1 μM GDC-0941 for 2 h and analyzed by western blot to detect total and phosphorylated levels of AKT. Actin is shown as the loading control. (Right panel) Viability of HCT116 D13/- pre-treated with 1 μM GDC-0941 for 2 h and then treated with 100 nM SM83 and 1 μM CPT. Cell viability was quantified after 24 h and expressed as viability percentage to inhibitor alone. (F) HCT116 cells were transfected with siRNAs targeting ERK1 and ERK2, cells were collected after 72 h and analyzed by western blot to detect Noxa and total and phosphorylated levels of AKT and ERK. Actin is shown as loading control.
in premalignant cells, but also in cancer cells. To this end, a panel of isogenic colon cancer cell lines with KI and KO mutations of KRAS was tested. In clear contrast to HME and MCF10A cells, Noxa levels did not depend on KRAS status in cancer cells and in line with this observation cell sensitivity was almost identical in each pair of isogenic cells. Basal levels of Noxa in cancer cells were higher than in epithelial cells (data not shown), suggesting that tumor cells constitutively express some pro-apoptotic proteins at high level, but could also activate parallel pro-survival pathways to counteract the pro-death signals supported by the MEK/ERK axis. Considerable evidence shows that mutations in the RAS/MEK/ERK cascade are associated to aberrant activation of PI3K/AKT signaling [37] and therefore both pathways should be targeted simultaneously for effective responsiveness to treatment [11, 38]. In accordance to this hypothesis, HCT116 cells and DLD1, which bear PI3K activating mutations, are sensitized to SM83/CPT treatment when pre-treated with AKT inhibitors only in the presence of oncogenic KRAS, supporting the notion that AKT is protecting from oncogenic KRAS-dependent cancer cell sensitization (Figure 7). It is important to note that this protective role was demonstrated also in cells bearing wild type PI3K (Figure S4), confirming the general pivotal role of AKT in counterbalancing the pro-death effect of oncogenic KRAS.

In conclusion, our work has two main implications. First, targeting down-stream effectors of oncogenes might result in an immediate and transient anti-proliferative effect often achieved by conventional therapies, but, more importantly, could also shut-down the pro-death signals derived from oncogene activation. Secondly, for a successful treatment, targeting of the EGFR/MAPK pathway alone is not sufficient [39], as it results in emerging protecting mutations [4, 40], feedbacks [41] and incomplete responses. It is therefore imperative to characterize and inhibit also the aberrantly activated survival pathways in a combination treatment, in order to overcome the anti-apoptotic effect of PI3K/AKT activation and simultaneously to exploit the pro-death signal stemming from oncogenic activation.

**Figure 7: Proposed mechanism for oncogenic KRAS-mediated sensitization to cell death.** In premalignant models, normal epithelial cells expressing endogenous or ectopic mutated KRAS express high levels of Noxa due to the hyper-activation of MAPK kinases and in particular of ERK2. In these settings, the basal activation of the PI3K/AKT pathway is not sufficient to protect from this pro-death stimulus and treatment with several cytotoxic agents results in Noxa-dependent cell killing. In contrast, in CRC cells, Noxa levels are independent of KRAS status and oncogenic KRAS-bearing cells respond to treatment to the same extent as in the presence of wild-type KRAS. In fact, mutated PI3K and up-stream stimuli likely deriving from the tyrosine kinase receptors activate AKT, which counterbalances the potential pro-death stimulus deriving from oncogenic KRAS.
METTHODS

Cell lines

The human isogenic hTERT-immortalized mammary epithelial cell lines HME +/+ and HME D13/+, and the human epithelial mammary MCF10A together with the isogenic pairs of colorectal cancer cell lines SW48 +/+ and SW48 D13/+, HCT116 +/+ and HCT116 D13/+, Lim1215 +/+ and Lim1215 D13/+, DLD1 D13/- and DLD1 +/- have already been described [4, 27]. HME isogenic pairs and MCF10A cell lines were cultured in DMEM-F12 (Gibco), supplemented with 10% FBS (LONZA), 2 mM glutamine (LONZA), 20 ng/ml EGF (Immunological Science), 10 µg/ml insulin (Sigma), 500 µg/ml hydrocortisone (Sigma-Aldrich). SW48 and DLD1 isogenic pairs were cultured with DMEM (Gibco) supplemented with 10% FBS and 2 mM glutamine. Lim1215 isogenic pairs were cultured with RPMI (LONZA) supplemented with 10% FBS, 2 mM glutamine and 1 µg/ml insulin. HCT116 were cultured in RPMI, supplemented with 2 mM glutamine, sodium pyruvate (LONZA), and non-essential amino acids (LONZA). HeLa cells for the high-throughput screening and packaging HEK293FT (Life Technologies) for lentiviral production were cultured in DMEM with 10% FBS. All cell lines were mycoplasma-free as determined by Takara Mycoplasma Detection Kit (Clontech).

Reagents

Antibodies targeting pan-RAS, Noxa (CalBiochem), Actin and ERK1/2 (Sigma), cleaved-PARP, cleaved caspase-3, phosphoERK1/2 (Thr202/Tyr204), pAKT and AKT (Cell Signaling), cIAP1 (R&D Systems), cIAP2 and XIAP (BD Biosciences), caspase-8 (Enzo Life Sciences) and Mcl-1 (Santa Cruz Biotechnology) were employed in western blot experiments. z-VAD(OMe)-FMK was purchased by BIOMOL, Necrostatin-1 from Enzo Life Sciences. PD98059 and UO126 were purchased from CalBiochem, LY294002 from Sigma, GDC-0941 and Triciribine from Selleckem. Infliximab (Schering-Plough) and Enbrel (Wyeth Pharmaceuticals) were used as TNF blockers. CPT and neocarzinostatin were purchased from Sigma-Aldrich, etoposide by Teva. SM83 synthesis has been described elsewhere [25, 28], while izTRAIL was purified as already shown [29]. Mutant KRAS (G13D) was cloned in the pMINDUCER20 and lentiviral particles prepared modifying an already described protocol [30] and using Lipofectamine 2000 as transfection reagent. Expression of the transgene was induced by doxycycline (Sigma-Aldrich) at the indicated concentrations.

High-throughput screening

On day 1, 350 HeLa cells/well were seeded in 384-well white plates in 20 µl medium. At day 2, media was changed with cells being exposed to 100 nM SM83 in addition to FDA-approved drug libraries (ENZO Life Sciences, MicroSource Discovery Systems Inc. and Prestwick Chemical, France) with the drug library compounds present at a final concentration of 1 µM. At day 3, cells exposed to SM83 were also exposed to 20 pg/ml izTRAIL. Cell viability was estimated on day 5 by CellTiter-Glo (Promega). Hits were selected due to their capability to enhance the cytotoxic activity of SM83/izTRAIL and then validated using the same HeLa cells employed in the screening. In validation experiments, SM83 and izTRAIL were administered alone and in combination, also changing the schedule and pre-treating cells with SM83 24 h before izTRAIL or administrating these compounds simultaneously. On the basis of these results, validated hits were tested in a panel of premalignant and cancer cell lines.

Western blot

Cells were harvested by centrifugation at 4500 rpm for 5 min at 4°C. After washing once with PBS, lysates were prepared by resuspending cell pellets in 60-100 µl lysis buffer (125 mM Tris HCl pH 6.8, 5 % SDS) supplemented with 1x complete protease (Roche Diagnostics) and phosphatase inhibitors (Sigma-Aldrich). Lysates were boiled at 99°C, sonicated for 20 seconds at RT. Then lysates were centrifuged at 13000 rpm for 20 min at RT and cleared supernatants were transferred into a new tube and frozen at -20°C. Cell lysates were mixed with 4x reducing SDS-Sample buffer containing 10 % β-mercaptoethanol (Sigma-Aldrich) and heated for 10 min at 99°C. Proteins (50 µg) were separated by SDS-PAGE using pre-cast 4-12% BisTris NuPAGE gels (Life Technologies), blotted to PVDF membranes (Millipore), which were washed with PBS-tween for 5 min, blocking buffer made of 4% non-fat milk in PBS-tween for 30 min and then incubated overnight with the indicated primary antibodies. Proteins were detected after hybridization with appropriate horseradish peroxidase (hrp)-conjugated secondary antibodies by adding a chemiluminescent substrate (EuroClone).

Cell viability assays

96-well optical bottom, polymer base white plates (Thermo Scientific) were used for viability tests. At day 1, 10000 cells per well were seeded in 100 µl medium. At day 2, cells were treated adding the indicated drug(s) in 10 µl volume per well. At day 3, cell viability was
determined using the CellTiter-Glo assay according to the manufacturer’s instructions. Statistical analysis was performed with GraphPad Prism 5.02 using the two-tailed unpaired t-test.

Transfection and lentiviral transduction

To achieve transient knock-down of target proteins in HME cells, a reverse transfection protocol employing siRNAs (Qiagen) and RNAiMAX (Life technologies) was used. Briefly, 3.25 µl RNAiMAX and 200 µl OptiMem (GIBCO) were mixed and incubated for 5 min at RT. Subsequently, 3.25 µl siRNA of a 20 µM stock were added, mixed and incubated for further 30 min at RT. The transfection mix was placed in a 6-well plate and 0.15 x 10^6 cells seeded in 800 µl on top.

Tumor cells were seeded the day before transfection and the same transfection mix as for reverse transfection was added on top of cells 24 h later. The cells were then incubated for 48 h or cultured for 72 h before stopping the experiment. In each experiment, scramble siRNAs (siCtr) were used as a control.

Cells transduced with lentiviral particles were cultured in the presence of medium collected from HEK293FT packaging cells transfected with pINDUCER20-KRAS G13D (referred to as G13D) or empty vector (Mock). After 48 h, medium was replaced and fresh medium added in the presence of 500 µg/ml G418 (Life Technologies).

Ras-GTP pull-down assay

2.5 x 10^6 cells were seeded into 10 cm dishes. The next day, cells were incubated with and without 250 ng/ml of Dox. Cell lysis and RAS-GTP pulldown was performed. Cells were lysed in 500 µl of IP-lysis buffer supplemented with a cocktail of protease inhibitors. To fully detach lysed cells, they were scratched using a cell scraper and transferred into tubes for a 30-minute lysis at 4 °C on a rotator. Lysates were centrifuged at 13000 rpm for 30 min and cleared supernatants were transferred to a new tube. RAS-GTP was precipitated using beads coated with the RAF1-binding domain RBD recombinant protein. The following day, beads were washed 5 times with IP-lysis buffer and precipitated protein complexes were eluted from the beads via boiling in SDS-Sample buffer for 10 minutes at 80°C. Proteins were separated by SDS-PAGE and analyzed by western blot. As a loading control, proteins were stained by blue coomassie (Thermo Scientific Pierce).

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