Cyclic AMP induces IPC leukemia cell apoptosis via CRE-and CDK-dependent Bim transcription

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The IPC-81 cell line is derived from the transplantable BNML model of acute myelogenic leukemia (AML), known to be a reliable predictor of the clinical efficiency of antileukemic agents, like the first-line AML anthracycline drug daunorubicin (DNR). We show here that cAMP acted synergistically with DNR to induce IPC cell death. The DNR-induced death differed from that induced by cAMP by (1) not involving Bim induction, (2) being abrogated by GSK3β inhibitors, (3) by being promoted by the HSP90/p23 antagonist geldanamycin and truncated p23 and (4) by being insensitive to the CRE binding protein (CREB) antagonist ICER and to cyclin-dependent protein kinase (CDK) inhibitors. In contrast, the apoptosis induced by cAMP correlated tightly with Bim protein expression. It was abrogated by Bim (BCL2L11) downregulation, whether achieved by the CREB antagonist ICER, by CDK inhibitors, by Bim-directed RNAi, or by protein synthesis inhibitor. The forced expression of BimL killed IPC-81WT cells rapidly, Bcl2-overexpressing cells being partially resistant. The pivotal role of CRE and CDK activity for Bim transcription is unprecedented. It is also noteworthy that newly developed cAMP analogs specifically activating PKA isozyme I (PKA-I) were able to induce IPC cell apoptosis. Our findings support the notion that AML cells may possess targetable death pathways not exploited by common anti-cancer agents.

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The second messenger cAMP regulates fundamental cellular processes, including cell survival, differentiation and proliferation, mainly by activating cAMP-dependent protein kinase I,II (PKA-I, PKA-II) or cAMP-stimulated exchange factors Epac1,2 (see1 and references therein). The IPC-81 acute myelogenic leukemia (AML) cells2 undergo apoptosis within 4–6 h after cAMP stimulation.3 The cells are also sensitive to first-line anti-leukemic drugs like daunorubicin (DNR) both in vitro and in vivo.4 This may reflect their origin from the BNML model of leukemia, considered reliable to judge the clinical efficacy of anti-leukemic drugs.5 The IPC cell provides therefore a unique opportunity to compare the cell death induced by a physiologically relevant endogenous second messenger and a chemotherapeutically relevant agent.

Intriguingly, the cAMP-induced IPC cell death is blocked by the cyclin-dependent protein kinase (CDK) inhibitor roscovitine (RCV),6 which (under trade name Seliciclib) is used against cancer.7,8 Equally intriguing, the cAMP-induced death is blocked by overexpression of ICER,9 which competitively inhibits the access of the cAMP-responsive gene promoter elements binding protein (CREB) transcription factor family members to CRE.10 ICER is considered an AML tumor suppressor and CREB an AML oncogene.11–13

The present study was undertaken to know how the normally proapoptotic RCV and ICER could block the cAMP-induced IPC cell death. We wanted also to know the relative roles of PKA-I, PKA-II and Epac in the death induction. For this purpose, new PKA-I selective cAMP analogs were developed and used, together with previously developed analogs discriminating between PKA and Epac.14,15 Finally, we wanted to know the relation, if any, between cAMP- and anthracycline-induced IPC cell death.

We found that activation of PKA-I was necessary and sufficient for cAMP-induced IPC cell death. The death was dependent on Bim induction, whose transcription was dependent both on CREB and RCV-inhibited CDK activity. In contrast, neither Bim induction, CREB activity or RCV-inhibitable CDK activity affected anthracycline-induced IPC cell death, which depended on glycogen synthase kinase 3β (GSK3β) activity and was promoted by disruption of the
HSP90/cochaperone p23 complex. In spite of having dissimilar apoptosis induction mechanism, cAMP synergized strongly with anthracycline to induce IPC cell death. We conclude that IPC cells have a potent cAMP-induced cell death pathway unexploited by first-line chemotherapeutics.

Results

Strong PKA-I activation induces IPC cell apoptosis, whereas the basal PKA activity is necessary to maintain growth and survival. The apoptosis induced (Figure 1a) by cAMP stimulation of IPC<sub>WT</sub> allow the probing of both cAMP signaling and apoptosis pathways. We studied first whether the cAMP receptor Epac could be co-responsible with PKA for the death induction. The Epac-specific activator 8-CPT-2'-O-Me-cAMP failed, however, to induce any apoptosis alone (Figure 1a) or to enhance the apoptosis induced by the PKA-specific activator N<sup>6</sup>-MB-cAMP (not shown; see also Supplementary Section IV). To probe the role of PKA more directly, the cells were stably transduced with short hairpin-based RNAi directed against the catalytic C<sub>a</sub> subunit of PKA, and selected for survival after 48 h exposure to an apoptogenic concentration of cAMP analog. The surviving clones expressed from 25–60% of the normal amount of catalytic kinase activity (Figure 1b). This suggests that at

![Figure 1](image-url)
least 60% of the normal PKA content is required for cAMP-induced apoptosis to occur. The RI subunit of PKA-I comprises about 75% of the total R subunit expressed in IPCWT cells, the remaining 25% being RII associated with PKA-II. An isolated activation of PKA-I might therefore be sufficient to induce apoptosis.

To enable specific activation of PKA-I, cAMP analog combinations with improved isozyme selectivity were synthesized. The new PKA-I selective analogs N6-Bnz-8-Pip-cAMP and 2-Cl-8-AHA-cAMP discriminate better than any previous cAMP analogs between the cAMP binding sites (A,B) of PKA-I and PKA-II (Supplementary Tables S1–S3). They acted in strong synergy to induce cAMP-dependent apoptosis (Figure 1c). When N6-Bnz-8-Pip-cAMP was combined with other cAMP analogs, a moderate (Figure 1d) or barely any (Figure 1e) synergy was achieved. The observed synergies are in accordance with predictions based on the analog cAMP binding site preferences (Supplementary Table S3), and therefore unlikely to be due to nonspecific side effects. Note also that the concentration of paired analogs yielding synergistic death induction was far lower than that required for each analog alone (Figure 1). We conclude that IPC cell apoptosis can be induced by the isolated activation of PKA-I.

Although a strong activation of PKA leads to IPC cell death, the basal activity seemed necessary for growth. In fact, some of the surviving shRNAi-transfected clones showed slow growth, sometimes associated with mitotic arrest. That the slow growth was due to low PKA activity was supported by the demonstration of improved growth of clone Lo45-II in the presence of a low concentration (20 μM) of cAMP analog (8-CPT-cAMP). This clone had about 25% of the normal PKA content against about 40–60% for three surviving clones that were unaffected by Bcl2 overexpression. The CDK inhibitor RCV nearly abolished the induction of Bim mRNA, indicating that Bim mRNA transcription involves a CDK-dependent step. The Bim mRNA expression appeared to be independent of new protein synthesis as it was unaffected by cycloheximide (CHX) (Figure 4a).

In conclusion, PKA appears to have a dual role. It induces apoptosis and affects mRNA expression via other transcription factors than CREB/CREM. The endogenous Bim expression is tightly coupled to death in cAMP-stimulated IPC cells. The CDK inhibitor RCV nearly abolished the induction of Bim mRNA, indicating that Bim mRNA transcription involves a CDK-dependent step. The Bim mRNA expression appeared to be independent of new protein synthesis as it was unaffected by cycloheximide (CHX) (Figure 4a).

As the cAMP-induced IPC cell death is blocked by both CHX and RCV, these findings suggested that Bim induction might be important for the death. If so, a correlation should exist between cell death and Bim protein expression, whether...
manipulated by interference with Bim-mRNA synthesis (RCV) or Bim-mRNA translation (CHX). We achieved a graded expression of Bim protein (see insets and right hand axes of Figures 4b and c) by adding RCV (Figure 4b) or CHX (Figure 4c) at successive time points after the addition of cAMP. In either case, Bim protein expression correlated tightly with apoptosis, as determined 6 h after the onset of cAMP stimulation.
To provide a more direct link between cAMP-induced apoptosis and Bim transcript induction, we knocked down the Bim transcript. The restriction enzyme-based method used to achieve IPC cells with knocked-down Cα (Figure 1b) was not feasible owing to lack of suitable restriction enzyme sites in the Bim-cDNA. We therefore designed synthetic RNAi hairpins to target the part of the Bim transcript coding for the common N-terminal part of all the detected Bim isoforms. The hairpin DNA was inserted into vectors to produce retrovirus and stably transduce IPCWT cells. Two of the selected clones were compared with a clone expressing non-target (luciferase) hairpin RNAi for apoptosis induction after incubation with cAMP analogs.

**Figure 2** Gene expression analysis of IPC cells exposed to the PKA-activating cAMP analog N6-MB-cAMP. IPCWT or IPC cells stably over-expressing ICER (IPCICER) or Bcl2 (IPCBCL2), were exposed for 2 h to the PKA activator N6-MB-cAMP (700 μM). The transcriptome was analyzed by Affymetrix gene array (U230A, 15,866 probe sets). (**a**) The 82 transcripts increased (red) or decreased (blue) >2-fold in either IPCWT or IPCICER or IPCBCL2 are shown in a hierarchical clustering diagram. Genes in green have known cAMP responsive elements (CRE) in the promoter region from −3000 to +300 bases from the transcriptional start site. (**b** and **c**) The Venn diagram circles show the number and overlap of transcripts upregulated (**b**) or downregulated (**c**) >2 fold by N6-MB-cAMP in IPCWT (grey), IPCICER (cyan), and IPCBCL2 (pink) cells. (**d**) The panel shows that the fold regulation by N6-MB-cAMP in IPCWT cells correlates well (r² = 0.866) with that in IPCBCL2 cells (Δ), but poorly (r² = 0.287) with that in the IPCICER cells (O). The dotted line represents perfect correlation. Note that only transcripts regulated more than 2-fold are included. (**e**) Biological processes over-represented among the N6-MB-cAMP regulated genes according to the http://www.pantherdb.org tool are shown. Two processes had significantly fewer and seven had more regulated genes than the average. The P-values ranged from 0.003 (intracellular protein transport) to 0.047 (protein metabolic process). The number of regulated genes for each process is shown above each bar (see Supplementary Table S5 for details).
8-CPT-cAMP. The hairpin-transduced cells were compared with non-target RNAi-transduced cells for ability to develop apoptosis in response to the cAMP analog 8-CPT-cAMP. The hairpin1 transduced cells showed very little apoptosis (Figures 5a and c) and low expression of Bim (Figure 5c, inset). Hairpin 2 gave less-efficient knockdown of Bim (Figure 5d) and protection against apoptosis (Figures 5a and d), but still provided significant protection as compared with the vector control (Figure 5b).

In conclusion, an excellent correlation is observed between Bim expression and apoptosis in cAMP-stimulated IPC cells, whether the Bim expression is manipulated by RCV, CHX or various Bim-directed shRNAi hairpins.

Enforced BimL expression induces IPC apoptosis independently of cAMP and more strongly in IPCWT than IPCBCL2 cells. In order to test whether Bim expression alone is sufficient to induce apoptosis in the IPCWT cells, they were transfected retrovirally with a bicistronic vector for BimL and GFP (BimL). Cells transfected with vector coding only for GFP served as control (pMIG). GFP-positive IPC WT and IPCBCL2 cells were observed 18 h after virus transduction with control vector. About 50% of the cells were detectably green 36 h after the transfection (not shown). No GFP positive cells were apparent after transfection with vector for with BimL + GFP. Nevertheless, a high proportion of the cells were apoptotic (Figure 5e). A likely explanation is that the retroviral-driven BimL expression induces apoptosis too rapidly for the cells to synthesize detectable GFP before undergoing death. The death morphology was classically apoptotic (Figure 5e, left inset), and strikingly similar to that induced by cAMP (Figure 1a).

The IPCBCL2 cells subjected to BimL + GFP transduction showed much less overall apoptosis than the similarly transduced IPCWT cells (Figure 5f). Some of the apoptotic cells were GFP positive (Figure 5f, left inset). This suggests that some of the BimL-transfected IPC BCL2 cells underwent apoptosis slowly enough to synthesize a detectable amount of GFP before dying. We conclude that ectopically expressed BimL is a highly efficient inducer of IPC cell apoptosis, whose effect is counteracted, but not completely abolished, by Bcl2 overexpression.

Cyclic AMP and DNR induce IPC cell death synergistically via distinct mechanisms. Cyclic AMP induces early mitochondrial damage in IPC cells. In spite of this similarity DNR does not affect Bim protein expression (Figure 6a). This may explain why DNR-induced death is independent on CDK activity (Supplementary Figure S1d) and insensitive to ICER overexpression (Supplementary Figure S3). Another distinction was that the effect of DNR, but not cAMP, was abrogated by inhibitors of the generally pro-apoptotic GSK3β protein kinase (Supplementary Figures S1b and c).

The ansamycin antibiotic geldanamycin (GA) antagonizes HSP90 by disrupting its association with p23. In spite of this similarity DNR does not affect Bim protein expression (Figure 6a). This may explain why DNR-induced death is independent on CDK activity (Supplementary Figure S1d) and insensitive to ICER overexpression (Supplementary Figure S3). Another distinction was that the effect of DNR, but not cAMP, was abrogated by inhibitors of the generally pro-apoptotic GSK3β protein kinase (Supplementary Figures S1b and c).

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lead to p23 truncation (Figure 6a). On the other hand, GA only enhances the DNR-induced IPC cell death (Figures 6b and e). This questioned the importance of the observed p23 cleavage for cAMP-induced death. To probe its importance, we prepared retroviral stable expression vectors for truncated p23 (Δp23), mimicking the endogenously formed p23 cleavage product, as well as a caspase cleavage-resistant (p23D142/145A) p23 (Figure 6d). We found no significant difference in cAMP-sensitivity between mock-transfected cells and cells expressing caspase-resistant p23 or Δp23 (Figure 6c). In contrast, the DNR-induced apoptosis was significantly higher in the Δp23-transduced cells than in the p23D142/145A-transduced cells (Figure 6f). That p23 cleavage can sensitize cells to DNR was supported further by experiments on HEK293T cells (Supplementary Figure S2).

We conclude that the cAMP- and DNR-induced cell death differs fundamentally regarding both Bim induction and effect of inhibition of CDK, GSK3β and HSP90/p23 activity. This suggests that DNR-resistant AML cells may have alternative death pathways that can be exploited therapeutically. In fact, cAMP and DNR synergized for IPC cell death induction (Figure 7). A considerable decrease of the cAMP analog dose required to induce apoptosis was achieved by using PKA-I synergistic cAMP analog pairs instead of single analogs (Figure 1). The use of analog pairs should therefore allow lessening of the cAMP analog load for potential clinical use. A further decrease should be achieved by acetoxy-methyl-ester prodrugs of the analogs, which have improved cell permeability.29

Discussion
Cyclic AMP signaling has versatile roles in hematopoiesis. They range from stimulation of proliferation to maturation and cell death.30,31 The presently studied IPC cell line reflects this versatility and places PKA as the pivotal cAMP mediator. PKA appears to have a delicate role in balancing the IPC cell proliferation. Cells expressing <40% of the C subunit of PKA
or being treated with PKA inhibitor (Rp-8-Br-cAMPS) are growth arrested. The high basal (resting state) cAMP level in IPC cells may therefore be an adaption required to activate the PKA enough to sustain proliferation. On the other hand, strong PKA activators arrest the cells at the G1/S restriction point.

Although the basal cAMP level is sufficient to sustain proliferation, it appears not to induce net CRE-dependent IPC cell transcription. Thus, the basal state IPC transcriptome is similar in wild-type cells and cells overexpressing the CRE-blocking ICER protein (not shown). In contrast, the gene expression is profoundly altered in ICER-transfected HL-60 cells. It is also puzzling in view of the notion of CREB as an AML oncogene and marker for therapy resistance and ICER as a tumor suppressor. The lack of CRE-dependent transcription in the basal state is not due to deficient CREB response. A number of transcripts are induced in an ICER-inhibitable manner in cells incubated with a PKA-directed cAMP analog. It may rather be related to the nature of the induced transcripts. They code for proteins involved in apoptosis or differentiation/development (Figure 1e), rather than growth and survival.

A near complete neutrophilic maturation can be achieved by cAMP stimulation of IPC cells stably overexpressing Bcl2.

**Figure 6** The cAMP-induced IPC cell apoptosis differs from DNR-induced apoptosis with respect to Bim expression and role of HSP 90 modulators. (a) Extracts from IPC cells treated with vehicle for 2 h (left Ctrl) or 6 h (right hand Ctrl), with 8-CPT-cAMP (200 μM) for 2 h, 4 h, or 6 h, or with 0.4 μM DNR for 6 h were probed for Bim, HSP90 co-chaperone p23 and actin by immunoblotting. Note that p23 cleavage is unaccompanied by Bim induction of the DNR-exposed cells. (b) Apoptosis was scored for IPC cells exposed to 0.5 μM GA alone or together with 8-CPT-cAMP (200 μM). (c) IPC cells retrovirally transfected with CRU5-IRES-GFP (empty vector), CRU5-IRES-GFP-p23 D142/145A (p23 D142/145A) or CRU5-IRES-GFP-p23 1–142 aa (Δp23) were treated with 8-CPT-cAMP (200 μM; 6 h) and scored for apoptosis. The 3–5% apoptosis in vehicle-treated cells is subtracted. (d) Cell lysates from cells transfected as for panel c were treated with vehicle (Ctrl) or DNR (0.4 μM; 6 h), and analyzed for p23, HSP90, and actin expression by immunoblotting. Note that cells transfected with Δp231–142 expressed Δp23 also before challenge with DNR. (e) IPC cells were treated with DNR (0.4 μM) in the presence or absence of GA (500 μM) and apoptosis scored. (f) The clones described in (c) were treated with DNR (0.4 μM; 6 h) and scored for apoptosis. Bars represent mean ± S.E.M., n = 3–8, and P-values are estimated using student t-test.

**Figure 7** Synergism between death induced by anthracycline and cAMP. IPCWT cells were preincubated for 2 h in the absence or presence of N6-MB-cAMP at a sub-apoptogenic concentration (175 μM) (Figure 1e). Thereafter the cells were added vehicle (0) or DNR to 0.1, 0.2, 0.3 or 0.4 μM concentrations. 3.5 h thereafter, the cells were fixed and assessed for apoptosis. Note the strikingly enhanced effect of DNR for cells pre/co-incubated with the cAMP analog. Data represent mean ± S.E.M. (n = 4).

Bcl2 can enhance maturation by directly affecting gene expression, as shown for the hematopoietic differentiation of murine embryonic stem cells. We find similar cAMP-induced transcriptomes of IPCWT and IPCBcl2 cells. Bcl2 may...
therefore promote differentiation indirectly, by simply allowing the cAMP-induced maturation program to progress without being overwhelmed by apoptosis, as in the IPCWT cells.

The most striking effect of IPCWT cell PKA activation is apoptosis induction, which occurs much more rapidly than in the S49 thymocyte cell line, which also depends on Bim induction. We found that Bim protein expression correlates closely with apoptosis. This is true whether Bim expression is manipulated by the CDK inhibitor (RCV), Bim-directed shRNAi, or by protein synthesis inhibitor (CHX). Bim is a plausible death inducer. It elicits intrinsic cell death by neutralizing the anti-apoptotic effects of Bcl2, Bcl-XL and Mcl-1, as well as through more direct pathways (see also Figure 8). In fact, the forced expression of Bim led to rapid death of IPCWT cells and more attenuated death of IPC BCL2 cells (Figure 5). A high proportion of the cAMP-induced Bim protein was of the BimEL variant, which is a substrate for PKA. It is therefore possible that cAMP may contribute to IPC cell death also by activating phosphorylation of the already induced BimEL protein. The role of Bim may reflect that the IPC cells have retained death pathway components operating in normal myeloid-derived cells. Bim is a negative regulator of normal hematopoiesis and accelerates the demise of mature neutrophilic granulocytes. It appears therefore that the IPC cells have retained normal bone marrow properties like pleiotropic response to cAMP and Bim-dependent death induction.

Figure 8 Overview of the pathway(s) incriminated in cAMP-induced IPC cell apoptosis. (1) The binding of cAMP to both sites on the regulatory subunits of the PKA holoenzyme leads to release of free, active catalytic subunits. (2) Catalytic PKA subunit translocated to the nucleus can phosphorylate the transcription regulatory proteins of the CREB family (CREB, CREM, ATFs), which bind to available CREs in the promoter region of CREB-regulated genes. The Bim-promoter contains four half CRE sites, two of which are conserved and separated by only 13 bp about 500 bp upstream from the transcription start site. The CREM isoform ICER blocks CRE sites and thereby the action of all CRE-binding transcription factors. Phosphorylated CREB recruits the basal transcriptional machinery via CREB-binding protein (CBP/P300) and other co-factors and adapters. (3) Transcript elongation is stimulated by phosphorylations of the C-terminal domain of RNA polymerase II by members of the CDK family, especially CDK7 and 9, which are sensitive to inhibition by RCV. (4) The translation of Bim-mRNA is inhibited by CHX. (5 and 6) The Bim protein can complex Bcl2 and thereby block Bcl2 prosurvival functions. Bim may (dotted line) also induce apoptosis by stimulating BAX/BAK directly. The dotted arrows from PKA to (5 and 6) is based on the observation that cAMP can synergize with DNR to induce apoptosis in IPCCEL cells (Supplementary Figure S3), which do not die in response to cAMP alone (Figure 1a). The dotted arrows from RCV to (5 and 6) are based on the moderate inhibition of cAMP-induced IPC cell death by dominant negative CDK5, which might be through inhibition of death processes downstream of Bim mRNA transcription.

The induction of Bim by cAMP has some unusual features. It occurs very rapidly, is mediated via CRE/CREB, and depends completely on CDK activity (Figures 2–4). The mediation by CRE/CREB is puzzling as the conserved CRE's of the Bim promoter are TATA-less and therefore not expected to be regulated by CREB. Antagonists of CDK7,9, like RCV, target the elongation of a subset of gene transcripts. The impact of CDK activity on Bim-expression appears not to have been explicitly studied, but is probably not universal. Thus, RCV causes depletion of anti-apoptotic proteins like Mcl-1 in neutrophilic granulocytes, but spares Bim, nevertheless our findings suggest some caution in using RCV in tumors depending on Bim expression for eradication.
The strong IPC cell response to anthracyclines is unsurprising in view of their origin from the transplantable BNML model of AML, considered a reliable predictor of the clinical efficacy of antileukemic drugs. The IPC cells are protected both against cAMP and the first-line anti-leukemic drug DNR. Furthermore, both death inducers promote the cleavage of the HSP90 co-chaperone p23 (Figure 6). In spite of these similarities, the cAMP and DNR-induced death did not proceed by a basically common pathway. Firstly, the p23 cleavage has consequence in DNR-induced death only (Figures 6b and e). Although the enforced expression of cleavage-resistant p23 protected and truncated p23 promoted DNR-induced death, they failed to affect cAMP-induced death (Figures 6c and f; Supplementary Figure S2). Similarly, the HSP90/p23-targeting drug GA only promoted the DNR-induced death (Figures 6b and e). Secondly, Bim is induced uniquely in cAMP-induced death (Figure 6a). Thirdly, CDK antagonism inhibits only the cAMP-induced death, whereas GSK3β antagonism only inhibits the DNR-induced death (Supplementary Figure S1).

In conclusion, the IPC cells demonstrate the co-existence of an extremely efficient PKA-CRE-CDK- and Bim-dependent pathway in addition to a p23- and GSK3β-enhanced anthracycline-induced death pathway. Studies aimed at exploring the therapeutic usefulness of the components of the PKA-CRE-CDK-Bim-dependent pathway may be warranted. It is of interest that PKA-I selective cAMP analogs can antagonize only in cAMP-induced death (Figure 6a). Thirdly, CDK antagonism inhibits only the cAMP-induced death, whereas GSK3β antagonism only inhibits the DNR-induced death (Supplementary Figure S1).

Materials and Methods

Reagents and constructs. The cAMP analogs were from BioLog (Bremen, Germany). The new A and B-site-specific CAMP analogs and the paullone analogs for CDK5 and GSK3β inhibition are described in Supplementary Section IV. RCY, GA and DNR were obtained from Sigma (St. Louis, MO, USA). The pMIG-Bim (Addgene plasmid 8786, Cambridge, MA, USA) was kindly made available for the scientific community by Dr. SJ Korsmeyer. The vectors for C2 knockout are described in Supplementary Section IV. To create expression vectors for shRNAi against Bim, HPLC-purified synthetic oligo DNA (see Supplementary Table S7b for sequences) were annealed and ligated into the Aar1 site of the retroviral shRNAi vector L087 (GenBank Acc: EU424173). Mutated and truncated versions of p23 were subcloned into vectors for transient transfection of HEK293T cells and stable retroviral transfection of IPC cells (see Supplementary Section IV for details).

Cell culture and retroviral transfection. Wild-type (cAMPWT), ICER- and Bcl2-overexpressing (IPCCER, IPCBCL2) clones of the IPC-81 leukemia cell line were obtained from Dr. M Lanotte, Hop St. Louis, Paris. They were grown in DMEM (Sigma) supplemented with 10% heat-inactivated horse serum (EuroClone, Milan, Italy), and kept in logarithmic growth phase. Before experiments, the cells were resuspended in fresh growth medium. The Phoenix retroviral producer cell line was a kind gift from Dr. J. Lorenz. They were grown in DMEM with 10% fetal bovine serum (EuroClone) and pen/strep as described above. By splitting every 2nd or 3rd day, confluent cells were avoided. The day before transfection, cells were plated at a density of 150 000 cells/cm². Transfection of phoenix cells was carried out by the calcium phosphate method. For a 10 cm² well, 2 μg VSV-G plasmid together with 5 μg plasmid of interest was used. Viruses were harvested after 24 (Bim expression) or 48 h (p23 and shRNAi expression), and purified by centrifugation at 50,000 g for 3 h, and resuspended in phosphate-buffered saline over night. IPC cells were infected in growth medium with protamine sulfate (5 mg/ml, Sigma) under centrifugation (1500 x g 90 min). For stable expression of Bim shRNAi, puromycin (Sigma) selection was started at 0.1 μg/ml 48 h after infection and gradually increased up to 1 μg/ml within 2 weeks.

Determination of apoptotic morphology and kinase activity. IPC cell apoptosis was assessed in formaldehyde-fixed (2% final concentration) cells. Apoptosis was assessed by phase contrast microscopy of the surface morphology, and fluorescence microscopy of the nuclear morphology after staining of DNA with bisbenzimide, Hoechst 33342 (Sigma). To determine cAMP-dependent protein kinase activity in IPC cells lysate, radioactive phosphate incorporation to kemptide was performed as previously described.29

Microarray analysis. IPC cell RNA was isolated with a combination of TRI reagent (Sigma) and GenElute columns (Sigma). The manufacturer’s protocol for TRI reagent was followed but instead of precipitation of the RNA, the aqueous solution was mixed with ethanol (final 35%) and applied to GenElute silica columns and washed and eluted according to the GenElute protocol. The microarray analysis on U230A rat transcriptome chip from Affymetrix (Santa Clara, CA, USA) was according to standard procedures at the Norwegian Microarray consortium Oslo. The processing of the expression data, annotations and cAMP responsive elements in the promoter region is described in the Supplementary Section IV.

Cloning of Bim transcript variants. RNA from 8-CPT-cAMP-stimulated IPC cells was reverse transcribed and amplified by PCR with primers given in Supplementary Table S7a. The PCR product was ligated into the pcDNA3/FRT/5′-His TOPO vector (Invitrogen, Carlsbad, CA, USA), and cloned to E.coli DH5α. Plasmid from ampicillin-resistant clones were isolated and sequenced. The sequences were aligned to known Bim transcripts and the genomic Bim sequence.

Quantitative RT-PCR and western blot. For quantitative RT-PCR, RNA was isolated with TRI reagent (Sigma) according to the protocol from the manufacturer. 1 μg of RNA was reverse transcribed with revert-aid Mu-MLV transcriptase (Fermentas GmbH, St. Leon-Rot, Germany) and oligo-dT primers and a specific reverse primer for 18S ribosomal RNA (18S). qPCR were run with a Lightcycler 480 II (Roche, Basel, Switzerland) using iQ SYBR green supermix (BioRad, Richmond, CA, USA). cDNA corresponding to 25 ng RNA were used in each PCR reaction. Primer sequences are given in Supplementary Table S7b. Bim expression was quantified relative to untreated control and the expression was normalized to the average of succinate dehydrogenase complex subunit alpha and 18S ribosomal RNA.

SDS gel electrophoresis and western blots were done and band intensity evaluated essentially as described previously.28 The membrane were probed with either: rabbit polyclonal anti-Bim (Cat No 202000, EMD Biosciences, San Diego, CA, USA), mouse monoclonal anti-flag (M2, Sigma), anti-Hsp90 (SPA-830, Stressgen, Victoria, Canada), mouse monoclonal anti-actin (ab6276 Abcam, Cambridge, UK) or mouse monoclonal anti-p23 (JJ3, generously supplied by Dr. David O Toft, Mayo Clinic, Rochester, MN, USA). Bands representing the proteins of interest were detected using alkaline-phosphatase-conjugated secondary antibodies (a-3687 and a-3562 Sigma) and CDP-Star substrate (Tropix, Bedford, MA, USA) and detected in a Fuji LAS3000.

Conflict of Interest

The authors S Huseby, G Gausdal, TJ Keen, C Krakstad, L Myhren, E Kjærland, K Brenstad, C Kunick, R Kleppe, and S Deskeland declare no conflict of interest. H-G Genieser is CEO and F Shvede is Head of Research and Development of BIOLOG Life Science Institute, which sells cyclic nucleotide analogs for research purposes.

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