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Experimental Hematology

Experimental Hematology 2018;00:00-00

Synergistic action of dual IGF1/R and MEK inhibition sensitizes childhood acute lymphoblastic leukemia (ALL) cells to cytotoxic agents and involves downregulation of STAT6 and PDAP1

Victoria J. Weston^a, Wenbin Wei^{a,b}, Tatjana Stankovic^a, and Pamela Kearns^a

^aInstitute of Cancer and Genomic Sciences, University of Birmingham, Edgbaston, Birmingham B15 2TT, UK; ^bSheffield Institute of Translational Neuroscience, University of Sheffield, Sheffield S10 2HQ, UK

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Heterogeneous upregulation of multiple prosurvival pathways underlies resistance to damage-induced apoptosis in acute lymphoblastic leukemia (ALL) cells despite normal p53 responses. Here, we show that the dual combination of insulin-like growth factor 1 (IGF1)/IGF1 receptor (IGF1/R) and mitogen-activated protein kinase/extracellular signal-regulated kinase (ERK) kinase (MEK) inhibition using AG1024+U0126 can sensitize apoptosis-resistant ALL cells to ionizing radiation-induced DNA damage irrespective of effect of single pathway inhibition in vitro. This AG1024 + U0126 combination also significantly potentiates the ability of the core chemotherapy compounds vincristine, dexamethasone, and daunorubicin to kill ALL cells in vitro. Evidence of the synergistic action of AG1024 + U0126 in samples with variable basal levels of phosphorylated IGF1/R β and ERK1/2 suggested additional targets of this drug combination. Consistent with this, gene expression profiling identified 32 "synergy genes" differentially targeted by IGF1/R + MEK inhibition and, among these, Signal transducer and activator of transcription 6 (STAT6) and platelet-derived growth factor-associated protein 1 (PDAP1) were the most differentially downregulated cluster. Pearson correlation analyses revealed that STAT6 and PDAP1 display significant expression codependency and a common expression pattern linked with other key "synergy" genes, supporting their predicted role in an STAT6–ERK–nuclear factor kappa beta (NF- κ B) network. Knockdown studies revealed that loss of STAT6, but not PDAP1, impinges on the cell cycle, causing reduced numbers of viable cells. In combination with daunorubicin, STAT6 loss has an additive effect on cell killing, whereas PDAP1 loss is synergistic, indicating an important role of PDAP1 in the cellular response to this anthracycline. Inhibition of STAT6 or PDAP1 may therefore represent a potential novel therapeutic strategy for resistant ALL by enhancing sensitivity to chemotherapy. © 2018 ISEH-Society for hematology and Stem Cells. Published by Elsevier Inc. This is an open access article under the CC BY license. (http://creativecommons.org/licenses/by/4.0/)

Although cure rates for childhood acute lymphoblastic leukemia (ALL) have increased dramatically over recent years through the integration of risk stratification into treatment protocols [1-3], failure of

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remission-inducing therapy is associated with only a 30% 10-year survival rate [4] and relapsed ALL remains the most common cause of cancer-related death in children [5-7]. There is a need for new therapeutic approaches with minimal toxicities for higherrisk leukemias.

Philadelphia-positive (Ph+) ALL is currently the only ALL subset for which treatment includes molecularly targeted treatment and the combination of imatinib with chemotherapy has significantly improved survival rates in this subgroup [8]. Recent advances in the molecular characterization of childhood ALL has

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Offprint requests to: Victoria J. Weston or Pamela Kearns, Institute of Cancer and Genomic Sciences, University of Birmingham, Vincent Drive, Edgbaston, Birmingham B15 2TT, UK; E-mail: victoria_weston@hotmail.com and p.r.kearns@bham.ac.uk

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identified several new, risk-associated genotypes implicating specific prosurvival signaling pathways [8–13]. Although these pathways potentially encompass multiple molecular targets, the evident heterogeneity could complicate a personalized treatment approach for ALL patients. For example, hyperactivation of the receptor tyrosine kinase and RAS signaling pathways is a prominent feature of hypodiploid ALL and these cells demonstrate sensitivity to phosphoinositide 3-kinase (PI3K), but not mitogen-activated protein kinase (MAPK)/extracellular signal-regulated kinase (ERK) kinase (MEK) inhibition in vitro [14]. In contrast, high-hyperdiploid ALL with activating RAS mutations is responsive to MEK inhibition in vitro [15]. Preclinical studies on adverse-risk Ph-like ALL, which frequently harbors Janus kinase (JAK) and cytokine receptor-like factor 2 (CRLF2) mutations, indicate sensitivity to JAK inhibition and mammalian target of rapamycin (mTOR)/PI3K inhibitors [10,16,17], whereas plateletderived growth factor beta (PDGFB)- and ABL-rearranged Ph-like disease instead appear to display responsiveness to the tyrosine kinase inhibitors imatinib and dasatinib [8,18,19]. Consistent with these findings, our previous work also showed that childhood ALL displaying both poor clinical outcome and impaired apoptotic responses to DNA damage in vitro exhibit heterogeneous upregulation of multiple prosurvival pathways, which notably involves the PI3K, insulin-like growth factor 1 (IGF1) and MAPK pathways [20,21]. Hyperactivation of these pathways, despite apparently normal p53 activation, causes sustained nuclear factor kappa beta (NF- κ B) activity and impaired apoptosis after DNA damage [20]. Treatment with single prosurvival pathway inhibitors could only sensitize ALL cells to DNA damage in vitro in a patient-specific manner [20,21].

Therefore, although it is clear that a significant proportion of ALL patients are likely to benefit from a molecularly targeted treatment added onto an existing chemotherapy regimen, the heterogeneous deregulation of signaling pathways paired with possible activation of compensatory pathways may hamper sensitization of ALL by single-pathway inhibition. We hypothesized that targeting dual pathways might be more effective against a broader spectrum of samples and indicate a more applicable therapeutic approach for patients with ALL showing impaired clinical responses. Here, we show that the specific combination of IGF1/IGF1 receptor (IGF1/R) and MEK inhibition can synergistically sensitize primary ALLs to a range of cytotoxic agents. We show that the mechanism of this drug combination involves downregulation of signal transducer and activator of transcription 6 (STAT6) and PDGFassociated protein 1 (PDAP1), which appear to function in a predicted STAT6-ERK-NF-*k*B regulatory network that may be implicated in apoptosis resistance in childhood ALL.

Methods

Patient ALL samples

Patient bone marrow (BM) samples were collected from Birmingham Children's Hospital with ethical approval and written consent (CCLG 08/H0405/22 and 08/H1208/4). Leukemic BM mononuclear cells were separated by density centrifugation and frozen in a viable state before use. For clinical data, see Supplementary Table E1 (online only, available at www. exphem.org).

Annexin V/propidium iodide (PI) apoptosis analysis

Cells were treated with 5 Gy ionizing radiation (IR) and incubated with U0126 (Promega, WI, USA), AG1024, or LY294002 (Calbiochem, Darmstadt, Germany) at 37°C for 24 hours, as described previously [21]. Apoptosis was assayed using an Annexin V Apoptosis Kit (BD Pharmingen, San Diego, CA) and analyzed using a Coulter Epics XL-MCL flow cytometer (Beckman Coulter, Fullerton, CA). Cells were considered to be apoptotic if they stained positive for both Annexin V and PI (Ann+/PI+). The proportion of IR-induced apoptotic cells was determined by subtracting the proportion of apoptotic cells detected in the absence of IR.

Drug preparation and cytotoxicity assays

U0126, AG1024, LY294002, vincristine, and daunorubicin were dissolved in dimethylsulfoxide (DMSO) and dexamethasone in 100% ethanol at stock concentrations of 10 mmol/L. Cells were incubated with U0126, AG1024, vincristine, daunorubicin, or dexamethasone (Sigma-Aldrich, St. Louis, MO, USA) at the indicated doses at 37°C for 72 hours and subsequently reacted with Cell TitreGlo reagent according to the manufacturer's instructions (Promega). Luminescence was quantified using a Victor Wallac plate reader.

Western blotting

Weston blotting was performed as described previously [21]. Antibodies included IGF1R β (#3027), phospho-IGF1R β (Tyr1131)/insulin receptor β (Tyr1146) (#3021), ERK1/2 (137F5), phospho-ERK1/2 (Thr202/Tyr204) (D13.14.4E), PDAP1 (#4300), procaspase 7 (D2Q3L), cleaved caspase 3 (5A1E) (Cell Signaling Technology, Danvers, MA, USA), STAT6 (ab44718) (Abcam, Cambridge, MA, USA), and PARP1 (Santa Cruz Biotechnology, Santa Cruz, CA, USA). Mouse monoclonal antibody against β -actin (AC-74) (Sigma-Aldrich) served as a loading control.

Microarray analysis

Cells were untreated, treated with 5 Gy IR, 30 μ mol/L AG1024+5 Gy IR, 20 μ mol/L U0126+5 Gy IR, or 30 μ mol/L AG1024+20 μ mol/L U0126+5 Gy IR for 6 hours in vitro before RNA extraction using a combined TRIzol/ chloroform(Invitrogen)/RNeasy column purification (Qiagen) method as described previously [21]. After first- and secondstrand synthesis and in vitro transcription, samples were hybridized to HuGene1.0 ST v1 gene chips (Affymetrix, Santa Clara, CA, USA). Probe-level quantile normalization [22] and robust multiarray analysis [23] on the raw .CEL files was performed using AltAnalyze [24]. Differentially

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expressed genes were identified using limma with a fold change >1.5 and p < 0.01 [25].

siRNA silencing of STAT6 and PDAP1

Small interfering RNA (siRNA) silencing was performed in HeLa cells using SiGenome SMARTpools targeting human *STAT6* (6778) and human *PDAP1* (11333) (Thermo Scientific, Waltham, MA, USA). Two daily sequential siRNA treatments were performed using DharmaFECT transfection (Thermo Scientific) according to the manufacturer's instructions and nontargeting pool #2 (scrambled siRNA) served as a control. Data from three separate knockdown experiments are shown.

Cell cycle analysis

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Treated cells were fixed in 100% ice-cold ethanol before staining with PI (Sigma-Aldrich) and cell cycle profiles assessed using a Coulter Epics XL-MCL flow cytometer (Beckman Coulter).

Statistical and network analysis

Combination indices (CIs) were determined using doseresponse curves and Calcusyn software and potentiation effects (PEs) using paired Student *t* tests of data normalized to untreated cells and with a single-agent effect subtracted. Standard deviations are shown. Pearson correlation coefficients were determined by comparing normalized LOG2 expression values as described previously [26]. Network analysis was performed by seeding synergy genes using Ingenuity Systems IPA software (Qiagen) according to the manufacturer's instructions.

Results

Combined IGF1/R and MEK inhibition using AG1024 + U0126 sensitizes primary ALL cells to DNA damage-induced apoptosis

As a consequence of the heterogeneous upregulation of multiple prosurvival signaling pathways underlying defective apoptotic responses in childhood ALL, we reported previously that individual prosurvival pathway inhibitors targeting MEK, IGF1/R, and PI3K induced patient-specific responses to IR-induced DNA damage in vitro [20,21]. We hypothesized that dual combination of prosurvival pathway inhibitors might sensitize a broader range of leukemias and therefore inform of a more applicable therapeutic approach. In this study, we compared dual combinations of the IGF1/R, MEK, and PI3K inhibitors AG1024, U0126, and LY294002, respectively, at the same micromolar doses we reported as single agents in our previous study [21]. We identified that the specific combination of AG1024 (IGF1/R inhibitor) and U0126 (MEK inhibitor) was consistently active in sensitizing four DNA damage-resistant ALL to IR, whereas dual combinations involving the PI3K inhibitor LY294002 induced more variable responses (Figures 1A and B). The combination of AG1024 + U0126 consistently induced the highest level of IR-induced apoptosis, as measured by Annexin and PI staining 24 hours after treatment (Figures 1A and B), and this occurred irrespective of the sensitivity of the cells to each single agent [21]. The drug vehicle DMSO alone had minimal effect on the survival of primary ALL cells at the same doses even after 72 hours (Supplementary Figure E1, online only, available at www.exphem.org). In two representative samples, western blotting confirmed that AG1024+U0126+IR induced caspase- and PARP1-dependent apoptosis to a greater extent than the single most potent inhibitor for each leukaemia. (Supplementary Figure E2, online only, available at www.exphem.org). Using doseresponse curves (Supplementary Figure E3, online only, available at www.exphem.org) and Calcusyn software, we were able to determine CI values for ALL-141, ALL-102, and ALL-106, which revealed strong synergism, synergism, and a near additive effect between AG1024 and U0126, respectively (Figure 1A and Supplementary Table E2, online only, available at www.exphem.org). For ALL-111, the dose curves for both U0126 and AG1024 alone were inhibitory. We therefore determined the PE that indicated a strongly synergistic effect of the combined AG1024+U0126 treatment (Figure 1B and Supplementary Figure E3 and Supplementary Table E2, online only, available at www.exphem.org). Therefore, whereas in ALL-141, ALL-102, and ALL-111, the combined effect of AG1024+U0126 was synergistic, in ALL-106, the combination of AG1024 + U0126 was not superior to U0126 alone: both single-agent treatments exhibited efficacy, as did the AG1024+LY294002 combination. Nonetheless, the sensitivity of ALL-106 to AG1024 and U0126 as single agents or in combination shows that a treatment strategy targeting IGF1/R + MEK pathways would sensitize apoptosis-resistant primary ALL cells to IR-induced DNA damage. We observed no synergistic effect of the two agents when combined in normal peripheral blood mononuclear cells in the presence of IR-induced DNA damage (Supplementary Figure E4, online only, available at www.exphem.org).

IGF1/R + *MEK inhibition using AG1024* + *U0126*

sensitizes childhood ALL to core chemotherapy agents We next investigated whether this specific combination of IGF1/R + MEK inhibition could sensitize ALL cells to other clinically relevant cytotoxic agents. To do so, we tested whether AG1024 + U0126 would sensitize ALL to the core remission-inducing chemotherapy agents dexamethasone, vincristine, and daunorubicin in vitro. We evaluated the effect of the lower doses 1 and 10 μ mol/L AG1024 + U0126 on chemotherapy-induced killing after 72 hours in eight primary ALL samples and observed chemosensitization for all three drugs (Figure 1C). In combination with 100 nmol/L dexamethasone, synergism was





observed with 1 and 10 μ mol/L AG1024 + U0126 (PE, paired *t* tests, *p* = 0.0084 and *p* = 0.0166, respectively). AG1024 + U0126 also increased ALL sensitivity to 100 nmol/L vincristine, with 1 μ mol/L displaying an additive effect and 10 μ mol/L displaying synergism (PE, paired *t* test, *p* = 0.019). Finally, AG1024 + U0126 exerted an additive effect with 10 nmol/L daunorubicin and, although synergism was observed with 100 nmol/L daunorubicin (paired *t* test, *p* = 0.01), the killing induced by daunorubicin alone was already very high and the incremental change was small (not shown). The combination of IGF1/ R + MEK inhibition at micromolar doses is therefore able to sensitize ALL cells, including high-risk and relapse samples, to submicromolar doses of chemotherapy agents in vitro.

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Variable basal levels of MEK and IGF1/R activity indicate additional mechanisms underlying the synergistic action of AG1024 + U0126

We next set about investigating the possible mode of action for the synergistic effect of AG1024+U0126 in sensitizing childhood ALL cells. To address this, we investigated whether an association existed between the basal levels of the phosphorylated ERK1/2, an indicator of activated MEK pathway, and phosphorylated IGF1/R β , an indicator of activated IGF1 and insulin pathways, in 10 primary ALLs that were sensitive to the synergistic effect of 10 μ mol/L AG1024+U0126 after 72 hours in vitro compared with each of the single inhibitors in the absence of other cytotoxic agents (PE, paired t test, p = 0.0001) (Figure 2A). We did not observe a synergistic effect using 1 μ mol/L AG1024 + U0126 (Supplementary Figure E5, online only, available at www.exphem.org). These samples (where material was sufficient) revealed highly variable basal levels of phosphorylated ERK1/2 and IGF1/R β proteins (Figure 2B), suggesting that the synergistic action of combined AG1024 + U0126 treatment was not wholly dependent on targeting the MEK and IGF1/R pathways.

Combined AG0124 + U0126 treatment induces a differential "synergy" gene profile

Based on the variable basal levels of activated of MEK and IGF1/R pathways, it appears that the synergistic

action of AG1024 + U0126 in ALL cells may function in part through additional or off-target activity. To explore possible additional mechanisms of this drug combination, we employed gene expression profiling in four ALLs, ALL-102, ALL-141, ALL-150, and ALLwhich displayed synergistic responses 200, AG1024 + U0126. Samples were treated either with IR to induce DNA damage, with each of the single inhibitors + IR, or AG1024 + U0126 + IR for 6 hours. To demonstrate that the individual inhibitors were active, we first evaluated the alteration of known target genes by comparing AG1024 + IR with IR alone and U0126+IR with IR alone. This comparison revealed differential expression of known target genes for each of the individual inhibitors, which, as well as MAPK G-protein-coupled receptor genes, and included miR222, SP1, and DUSP6 [27,28] for the MEK inhibitor U0126 and JUN, SPRY, APAF1, and BAK1 [29] for the IGF1/R inhibitor AG1024, as well as TNS3, previously correlated with IGF1 levels [30]. We subsequently compared gene expression in cells treated with AG1024 + U0126 + IR with cells treated with each inhibitor alone + IR to identify a set of "synergism" genes that were differentially expressed after 6 hours of treatment with AG1024 + U0126 (for Venn analysis, see Supplementary Figure E6, online only, available at www.exphem.org). Of the 32 differentially expressed genes, six were consistently differentially upregulated and 26 downregulated by AG1024+U0126+IR treatment (Figure 2C). Because this set of genes could contribute to apoptosis resistance in some ALL, we set about investigating candidate synergy genes further.

Synergy genes STAT6 and PDAP1 exhibit codependent expression

Of the set of synergy genes we identified, *STAT6* and *PDAP1* were among the most significantly differentially downregulated (Figure 3A) and were validated by independent quantitative reverse transcription polymerase chain reaction (Supplementary Figure E7, online only, available at www.exphem.org). These genes were of particular interest because of their possible role in high-risk subtypes of childhood ALL.

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each treatment is shown, as well as the average IR-induced percentage of apoptosis (shown in parentheses), which were determined by subtracting the background apoptosis from the IR plots. (**B**) The combination of AG1024+U0126 (shaded) consistently sensitizes ALL cells to IRinduced apoptosis regardless of the effect of AG1024 and U0126 as single agents in four cases (ALL-111, ALL-102, ALL-106, and ALL-141) after 24 hours of treatment, as determined by Annexin V/PI staining. (The maximum concentration of DMSO reached in this experiment was 0.6% v/v DMSO.) CIs could be determined from individual drug dose curves (Supplementary Figure E3, online only, available at www.exphem. org) for ALL-102 (++, moderate synergism), ALL-106 (+, nearly additive), and ALL-141 (+++, synergism). For ALL-111, the PE of the combination of AG1024+U0126 was determined using unpaired t test analysis as described (***, strong synergism) (Supplementary Table E2, online only, available at www.exphem.org). (**C**) AG1024 and U0126 together sensitize ALL (S025117, S032957, S026767, ALL-202, ALL-201R, ALL-211, ALL-203, and ALL-212) to 100 nmol/L dexamethasone (top) 100 nmol/L vincristine (middle), and 10 nmol/L daunorubicin (bottom) after 72 hours of treatment. (The maximum DMSO concentration reached was 0.2% v/v DMSO.) The PE of AG1024+U0126 with each drug was determined using paired t test analysis. ** $p \le 0.005$ indicates moderate synergism and * $p \le 0.05$ indicates synergism.





Figure 2. The synergistic effect of combined inhibition of IGF1/R + MEK pathways by AG1024 + U0126 induces a distinct "synergy" gene signature. (A) In the absence of cytotoxic agents, primary ALL samples (n = 10) display synergistic sensitivity (PE = ***) to treatment with both

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Phosphorylated STAT6 has been reported to be elevated in Ph+ ALL [31] and PDAP1 appears to be involved in PDGF signaling [32,33], a pathway recently implicated in apoptosis-resistant ALL [21] and Ph-like ALL [8-12,18]. To provide some insight into how these synergy genes might interact to induce chemosensitization when downregulated, we used logarithmic expression data from 20 arrays used in this study (untreated, IR only, AG1024+IR, U0126+IR, and AG1024+U0126+IR for each of the four ALL samples) to enable Pearson coefficient correlation analyses. This identified a marked (r > 0.6) and highly significant expression codependency between STAT6 and PDAP1 (Figure 3B). When we compared a possible relationship between STAT6 and PDAP1 and the remaining synergy genes, we observed a striking pattern of linked expression that was common to both STAT6 and PDAP1 (Figure 3C).

Identification of a predicted regulatory network underlying the synergistic activity of AG1024 + U0126 in ALL cells

We next performed network analysis using Ingenuity software to determine whether the synergy genes that we identified were predicted or known to interact in common pathways. This analysis identified a major network associated with cell death and survival, cellular function and maintenance, and hematological system development and function. STAT6 and PDAP1 were associated with this network, of which STAT6 was a major component in addition to ERK and NF- κ B. In this STAT6–ERK–NF- κ B network, STAT6 and PDAP1 appeared to be closely associated (via PDGF), consistent with our codependent expression data (Figure 3D). Other components of this network (PRDM8 and MPZL) had also shown linked expression with both STAT6 and PDAP1 using Pearson correlation analysis (see above). This STAT6-ERK-NF-KB network could be an important mechanism underpinning apoptosis resistance in ALL cells and warrants further characterization.

Knockdown of STAT6 and PDAP1 has an impact on cell cycle and chemosensitization

Because our data suggested that STAT6 and PDAP1 might contribute to apoptosis resistance, we hypothesized that loss of these proteins individually might reduce cell viability and/or sensitize cells to chemotherapy. To address the cellular impact of loss of STAT6 and PDAP1, we performed siRNA-mediated knockdown in HeLa cells. Because of the difficulties in obtaining gene knockdown in primary human ALL cells, we used HeLa cells as an alternative since they enable reproducible and effective protein knockdown. Although HeLa cells are of a different tissue origin than ALL, the rationale for choosing this model system to explore the cellular impact of loss of these proteins on chemotherapy sensitization was because they also represent a widely accepted model for studying DNA damage response proteins [34,35]. Figure 4A shows reduction of each of the proteins after 72 hours of siRNA treatment. Interestingly, we observed that the loss of PDAP1 appeared to lead to an increase in STAT6 protein expression. This apparent connection at the protein level supports the codependent expression correlations that we observed at the mRNA level for these molecules.

We first investigated the impact of loss of each of the proteins on cell viability and the cell cycle. In untreated cells, STAT6 knockdown by siRNA treatment had the greatest impact on the number of viable cells, whereas PDAP1 loss had no impact on the number of viable cells after 1 week of siRNA treatment (Figure 4B). When we assessed alterations in the cell cycle, we found that, consistent with the reduction in the number of viable cells and with previous findings [36], STAT6 caused a 20% increase in the proportion of cells in the G_1 phase of the cell cycle, indicating an increased number of arrested cells and a reduction in cycling. In contrast, PDAP1 loss had no impact on the cell cycle, which is consistent with the absence of effect of PDAP1 loss on viable cell number (Figure 4C).

We next investigated the ability of loss of each of these proteins to sensitize cells to core chemotherapy agents used to induce remission in ALL patients. When we combined protein knockdown with increasing doses of the anthracycline daunorubicin in vitro, the impact on the total number of viable cells caused by knockdown of STAT6 had an additive effect (Figure 4D), whereas, strikingly, PDAP1 knockdown was synergistic after 72 hours (Figure 4D). In contrast, when STAT6 was silenced, the impact on cell cycle arrest (Figure 4D) reduced the sensitivity of HeLa cells to increasing doses of vincristine, the mechanism of which is dependent on cell proliferation. PDAP1

AG0124 + U0126 after 72 hours. (The maximum DMSO concentration reached was 0.2% v/v DMSO.) (**B**) Western blot analysis revealing samples displaying synergism to AG1024 + U0126 and variable basal MEK and IGF1/R pathway activity, as indicated by phosphorylation of IGF1/ R β + IR β and ERK1/2, suggesting that additional mechanisms underlie the synergistic effect. (**C**) Heat map showing differential expression of 32 genes ("synergy" signature) induced by 6 hours of treatment with AG1024 + U0126 compared with the respective individual inhibitors + IR, IR alone, and untreated cells in ALL-102, ALL-141, ALL-150, and ALL-200. * $p \le 0.05$ indicates synergism; *** $p \le 0.0005$ indicates strong synergism.

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Figure 3. Predicted interactions of "synergy" genes *STAT6* and *PDAP1*. (A) Heat map showing that *STAT6* and *PDAP1* are highly differentially downregulated by AG1204 + U0126. (B) Pearson correlation coefficient analyses from 20 microarrays indicating that *STAT6* and *PDAP1* expression are highly codependent. (C) *STAT6* and *PDAP1* display common patterns of expression c-dependency with specific "synergy" genes highlighted in bold text. (Pearson coefficient *r* values: below 0.4 = weak; 0.4 - 0.6 = modest; above 0.6 = marked codependency. (D) Ingenuity network analysis predicting interaction of *PDAP1* and *STAT6* in a STAT6–ERK–NF- κ B regulatory network seeded with "synergy" genes that might be involved in apoptosis resistance (green = downregulated; red = upregulated). * $p \le 0.05$; ** $p \le 0.0005$; *** $p \le 0.0005$.



Figure 4. Cellular consequences of siRNA-mediated knockdown of synergy genes. (A) Western blot showing siRNA mediated knockdown of STAT6 and PDAP1 in HeLa cells. (B) siRNA-mediated knockdown of STAT6 results in a significant loss in cell viability, whereas PDAP1 knockdown has no impact after 1 week. (C) siRNA-mediated knockdown of STAT6 causes a 20% increase in cells in the G₁ phase of the cell cycle, indicating decreased proliferation, whereas PDAP1 knockdown has no impact on the cell cycle in HeLa cells. (D) Effect on loss of cell viability by STAT6 knockdown leads to an additive effect with daunorubicin after 72 hours of treatment, whereas PDAP1 knockdown is synergistic with daunorubicin at more than one dose in HeLa cells. * $p \le 0.05$; ** $p \le 0.005$.

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knockdown had no effect at all on vincristine-induced killing after 72 hours cell of treatment (Supplementary Figure E8, online only, available at www.exphem.org). The HeLa cells displayed no evidence of sensitivity to dexamethasone with or without knockdown (data not shown). These data support the notion that the inhibition of pathways involving STAT6 and PDAP1 contributes to the potentiating effect of combined AG1024+U0126 treatment on DNA-damaging agents such as IR and daunorubicin. The inability of loss of the single genes selected for further exploration in this study to sensitize HeLa cells to vincristine or dexamethasone suggests that it is likely that alternative genes or a combination of genes are implicated in sensitization observed in both HeLa and leukemia cells. This is supported by the microarray data. However, with respect to daunorubicin, our data from HeLa cells indicate that STAT6, and in particular PDAP1, appear to function in the cellular response to daunorubicin and warrant further investigation in leukemia cells.

In summary, we have identified a specific combination of prosurvival signaling pathway inhibitors, AG1024 + U0126, which was consistently able to sensitize apoptosis-resistant primary ALL cells to DNAdamaging agents. This dual combination targets a specific set of "synergy" genes that includes *STAT6* and *PDAP1*, which are predicted to function in an STAT6-ERK-NF- κ B regulatory network.

Discussion

Childhood ALL is genetically heterogeneous and deregulation of different prosurvival signaling pathways can contribute to apoptosis resistance. In this study, we sought to determine whether the inhibition of a specific combination of prosurvival pathways could sensitize ALL irrespective of response to single pathway inhibition and if this might inform of a more applicable uniform treatment approach for ALL patients. We have shown that the combined inhibition of the IGF1/R and MEK pathways using AG1024 + U0126 can sensitize ALL cells in a synergistic manner with IR-induced DNA damage and can also potentiate the effects of core chemotherapy agents in vitro. Gene expression profiling revealed a set of synergy genes that included STAT6 and PDAP1, which are predicted to function in a STAT6-ERK-NF-kB regulatory network.

JAK/STAT signaling is implicated in many cancers [36]. STAT6 hyperactivation has been described in a several lymphoid malignancies and, recently, STAT6-activating mutations have been identified in a range of lymphomas with a frequency of up to 40% [36–39]. In ALL, phosphorylated STAT6 levels are elevated in Ph+ disease [31] and, given the similarities, it is interesting to speculate that STAT6 activity might also be

upregulated in the high-risk Ph-like ALL subtype. STAT6 signaling has been implicated specifically in treatment resistance and progression in several malignancies, including the response of chronic lymphocytic leukemia cells to B-cell receptor-mediated treatment and radioresistance in inflammatory breast cancer cells [40-43]. There is mounting evidence, therefore, that STAT6 signaling plays a relevant role in the pathophysiology and clinical responses of lymphoid and other malignancies. These data support our conclusions that STAT6 is likely to be important in the cellular response to DNA damage and could contribute to apoptosis resistance in ALL.

Although the function is largely unknown, PDAP1 was originally identified as a PDGF-interacting protein [32]. Indeed, PDAP1 was shown recently to be an effector of PDGR signaling in glioma cells and was associated with proliferation and disease progression, highlighting PDAP1 as a potential therapeutic target [33]. PDGF signaling is a pathway that we recently implicated in apoptosis resistance in childhood ALL [21] and, furthermore, *PDGFRB* translocations are a recurrent feature of high-risk ALL [8-12,18]. Interestingly, cells with NF1 inactivation and consequential hyperactivation of the RAS-MAPK-ERK signaling pathway also display overexpression of PDGFRs, which contributes to RAS-induced proliferation [44,45] and supports a role for PDAP1 in a putative STAT6-ERK-NF-kB regulatory network. In our study, we showed that loss of PDAP1 synergized significantly with daunorubicin to induce killing in HeLa cells. We suggest that the role of PDAP1 in the cellular response to DNA-damaging agents should be investigated further in leukemia cells because PDAP1 might represent an interesting novel therapeutic target for chemosensitization.

In summary, through dual IGF1/R and MEK inhibition, we have identified a group of genes that appear to contribute to impaired apoptotic responses to DNA damage and, when targeted, can sensitize ALL cells to chemotherapy agents. In particular, we have demonstrated that STAT6 and PDAP1, via a putative STAT6-ERK-NF- κ B network, may represent useful molecular targets for treatment-resistant ALL, particularly in the absence of clinically available IGF1/R inhibitors. It has already been demonstrated that JAK inhibitors such as ruxolitinib or leflunomide, which abrogate JAK3/STAT6 tyrosine phosphorylation, could represent useful treatment approaches for some ALL. A novel therapeutic approach for ALL could also potentially include STAT6 inhibition. STAT6 smallmolecule inhibitors are currently under development for the treatment of asthma and could also be evaluated in the context of ALL. Overall, it will be important to delineate STAT6/PDAP1 signaling precisely in ALL

and other malignancies to elucidate the role of these molecules in treatment resistance.

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Conflict of interest disclosure

The authors declare no competing financial interests.

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	able E1. Clinical features	of paediatric ALL sam	ples.	WCC (50-10 ⁹ /L)	
R025117	sall	Age (y)	ETV4 DUNV1	108	
S025117 S026767	CALL	5.1 7.08	EIVO-KUNAI	108	nk
5020707	I D	7.08		121.4	
5027830	D D	2.7		13.5	
5029946	В	11	IgH@	10.1	HK
5029947	В	2	High hyperdiploid	9.3	HK
5032957	В	5.11	High hyperdiploid	10.2	нк
5038556	в	9.9	High hyperdiploid	33.8	
ALL-75	Т	7	2xp16del	193	nk
ALL-102	cALL	7.02	Hyperdiploid (52)	62	HR
ALL-106	cALL	15	Near Haploid (28)	117	HR
ALL-111	cALL	10.09	Gain of AML1	2	HR
ALL-115	Т	4.1	Myb ^{aup}	nk	LR
ALL-141	cALL	14	Gain of ETV6	11	HR
ALL-150	В	8	ETV6-RUNX1	nk	LR
ALL-200	В	4.06	nk	142	nk
ALL-201R	BCP-ALL	17.5	IGH@	na	na
ALL-202	cALL	6.08	46,XX Subclone of IGH@ gain	3	LR
ALL-203	cALL	2.10	CRLF2 rearranged	17.6	HR
ALL-211	Т	7.05	TCRD-LM02	405	LR
	т	5.07	46, XY	140	LR
ALL-212	1		SIL-TAL1	493	HR
ALL-212 ALL-213 Nk, not known	; na, not applicable, F	8 IR, High risk; LR,	Low risk.	775	
ALL-212 ALL-213 Nk, not known Supplementary T IR.	T ; na, not applicable, F able E2. Combined effect	8 IR, High risk; LR, of U0126+AG1024 in	Low risk.	473	
ALL-212 ALL-213 Nk, not known Supplementary T IR. ALL	able E2. Combined effect	8 IR, High risk; LR, of U0126+AG1024 in of U0126+AG1024	Low risk.		
ALL-212 ALL-213 Nk, not known Supplementary T IR. ALL ALL-141 ALL-142	T ; na, not applicable, F able E2. Combined effect Combined effect	8 IR, High risk; LR, of U0126+AG1024 in of U0126+AG1024 +++ synergi:	Low risk.		
ALL-212 ALL-213 Nk, not known Supplementary T IR. ALL ALL-141 ALL-102 ALL-102	T T ; na, not applicable, F able E2. Combined effect Combined effect CI=0.478 CI=0.714 CL = 0.22	8 IR, High risk; LR, of U0126+AG1024 in of U0126+AG1024 +++ synergi: ++ moderate	Low risk. ALL treated with		
ALL-212 ALL-213 Nk, not known Supplementary T IR. ALL ALL-141 ALL-102 ALL-106 ALL-111	T T ; na, not applicable, F able E2. Combined effect Combined effect CI=0.478 CI=0.714 CI=1.020 PE = -0.0001	8 IR, High risk; LR, of U0126+AG1024 in of U0126+AG1024 +++ synergi: ++ moderate + nearly add	Low risk. ALL treated with sm synergism itive		

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72h (n=5) 24h (n=5) 2. Surviving fraction of cells Surviving fraction of cells 0.5 0.5 0.25 0.25 0.125 0.125 0.2 0.5 0.6 0.2 0.5 0,6 Percentage DMSO (v/v) Percentage DMSO (v/v) Supplementary Figure E1. Graphs show minimal effect on survival of cells from 5 representative primary ALL samples following 24h and 72h treatment with doses of DMSO (drug vehicle) reflecting those doses reached within the experimental data (<0.6% DMSO). In contrast, 1% DMSO did cause a cytotoxic effect after 72h.

		ALL-111			ALL-141			
LY294002 (30µM):	-	-	+	-	-	-	-	-
U0126(20µM):	-	-	-	+	-	-	-	+
AG1024(30µM):	-	-	-	+	-	-	+	+
IR (5Gy):	-	+	+	+	-	+	+	+
PARP1 cleaved PARP1	-	-	-	-		-	=	11
procaspase 7	-	-	-	-	-	-	-	
cleaved caspase 3			-	=	-	-	=	=
actin	-	-	-	-	-	-	-	-

Supplementary Figure E2. Western blot analysis comparing caspase-dependent apoptosis induced by AG1024+U0126 with the single most potent single inhibitor 8h following treatment in two representative ALL (ALL-111 and ALL-141). In ALL-111, which exhibited equal sensitivity to LY294002 and U0126+AG1024, caspase and PARP cleavage were comparable following 8h treatment. In the completely IR-resistant leukaemia, ALL-141, caspase and PARP1 cleavage was significantly induced by treatment with U0126+AG1024 compared with AG1024 alone, the only single inhibitor with some effect.

12.e2

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Supplementary Figure E4. Graph shows absence of a combined effect of AG1024+U1026 on IR-induced apoptosis in peripheral blood mononuclear cells (PBMCs) from three healthy individuals, measured by Annexin V/PI staining and FACS analysis after 72h.



Supplementary Figure E5. Graph shows no synergism between AG1024 and U0126 at a dose of 1μ M following 72h treatment in 10 primary ALL.

12.e3

1604	12.e4	V	. Weston et al. / Experimenta	l Hematology 2018;00:00-00		1661
1606		Differentially days	n rogulated gapage	Differentially up r	agulated gapage	1663
1607		Differentially dow	n-regulated genes.	Differentially up-i	egulated genes.	1664
1608		AG1024+U0126+IR	AG1025+U0126+IR	AG1024+U0126+IR	AG1025+U0126+IR	1665
1609		VS	VS	VS	VS	1666
1610		AG1024+IR	00126+IR	AG1024+IR	U0126+IR	1667
1611			\frown			1668
1612						1669
1613						1670
1614		140 24	176	155 5	72	1671
1615						1672
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1620		•	,		/	1679
1622						1679
1622						1680
1623		PET1	12	GC	M2	1681
1625		REXA	NK. 24	GALI	NT14	1682
1626			D1	AC012	v	1683
1627		ACY	P1	C11c	r rf41	1684
1628		PAL	LD	0110		1685
1629		CAR	D6			1686
1630		TNS	3			1687
1631		ALP	(3			1688
1632		CAS	P1			1689
1633		PRU	NE			1690
1634		KCN	NB			1691
1635		PRDI	V18			1692
1636		DNA	H3			1693
1637		GLD1	.L.3 T.6			1694
1638		ARN ARN	T2			1695
1639		SUM	03			1696
1640		CRYM	AS1			1697
1641		S100	A2			1698
1642		MPZ	L1			1099
1644		Ue	5			1700
1645		CARE)16			1701
1646		MIR2	222			1702
1647	Supplementar	v Figure E6. Identification of	f genes contributing to AG10	24+U0126 synergism. Venn	analysis identifying 'synergy' genes shows	1704
1648	26 genes are	differentially down-regulated	(left) and 6 genes are diffe	rentially upregulated (right)	by treatment of ALL cells with AG1024	1705
1649	+U0126+IR co	mpared with AG1024+IR and	U0126+IR in samples ALL-	102, ALL-141, ALL-150 and	ALL-200.	1706
1650						1707
1651						1708
1652						1709
1653						1710
1654						1711
1655						1712
1656						1713



no effect compared with scrambled siRNA, following 72h treatment in vitro.