Investigating chemoresistance to improve sensitivity of childhood T-cell acute lymphoblastic leukemia to parthenolide

Benjamin C. Ede,1 Rafal R Asmaro,1 John P. Moppett,1,2 Paraskevi Diamanti1,3 and Allison Blair1,3
1School of Cellular and Molecular Medicine, University of Bristol; 2Bristol Royal Hospital for Children and 3Bristol Institute for Transfusion Sciences, NHS Blood and Transplant, UK

ABSTRACT

Current therapies for childhood T-cell acute lymphoblastic leukemia have increased survival rates to above 85% in developed countries. Unfortunately, some patients fail to respond to therapy and many suffer from serious side effects, highlighting the need to investigate other agents to treat this disease. Parthenolide, a nuclear factor kappa (κ)B inhibitor and reactive oxygen species inducer, has been shown to have excellent anti-cancer activity in pediatric leukemia xenografts, with minimal effects on normal hemopoietic cells. However, some leukemia initiating cell populations remain resistant to parthenolide. This study examined mechanisms for this resistance, including protective effects conferred by bone marrow stromal components. T-cell acute leukemia cells co-cultured with mesenchymal stem cells demonstrated significantly enhanced survival against parthenolide (73±11%) compared to cells treated without mesenchymal stem cell support (11±9%). Direct cell contact between mesenchymal cells and leukemia cells was not required to afford protection from parthenolide. Mesenchymal stem cells released thiols and protected leukemia cells from reactive oxygen species stress, which is associated with parthenolide cytotoxicity. Blocking cystine uptake by mesenchymal stem cells, using a small molecule inhibitor, prevented thiol release and significantly reduced leukemia cell resistance to parthenolide. These data indicate it may be possible to achieve greater toxicity to childhood T-cell acute lymphoblastic leukemia by combining parthenolide with inhibitors of cystine uptake.

Introduction

The introduction of contemporary therapies for childhood T-cell acute lymphoblastic leukemia (T-ALL) has resulted in remission rates that are closer to that of B-cell precursor (BCP) ALL but survival rates remain lower and 15-20% of children with T-ALL die from relapsed/refractory disease.1 Patients with high-risk disease or those who relapse often receive more intensive treatment, making them more susceptible to toxicity and long-term secondary complications.2 This highlights the need to investigate other agents to treat this disease.

It has been demonstrated that numerous cancers generate high levels of reactive oxygen species (ROS) compared to healthy tissue counterparts, where ROS levels are normally maintained in a tightly controlled manner.3 In T-ALL, ROS levels have been shown to be heightened, and this can inactivate phosphatase and the tensin homolog (PTEN), promoting leukemia cell survival.4 In human T-ALL, ROS levels are restrained by downregulation of protein kinase c theta (PKCθ) caused by NOTCH-1, a commonly activated mutation in T-ALL.5 However, if ROS stress levels are pushed above a certain threshold, cell death is forced to occur.6 Therefore, ROS promoting drugs may be an effective way of targeting cancer cells. Parthenolide (PTL) has been previously shown by ourselves and others to be a promising therapeutic agent for blood cancers.6-8 Importantly, it has limited effects on normal cells.

Correspondence:
allison.blair@bristol.ac.uk

Received: December 18, 2017.
Accepted: May 10, 2018.
Pre-published: May 17, 2018.

Check the online version for the most updated information on this article, online supplements, and information on authorship & disclosures: www.haematologica.org/content/103/9/1493

©2018 Ferrata Storti Foundation
Material published in Haematologica is covered by copyright. All rights are reserved to the Ferrata Storti Foundation. Use of published material is allowed under the following terms and conditions:
https://creativecommons.org/licenses/by-nc/4.0/legalcode. Copies of published material are allowed for personal or internal use. Sharing published material for non-commercial purposes is subject to the following conditions:
https://creativecommons.org/licenses/by-nc/4.0/legalcode, sect. 3. Reproducing and sharing published material for commercial purposes is not allowed without permission in writing from the publisher.
on normal cells at the doses required to kill cancer cells. PTL can target cancer cells via numerous mechanisms, such as inhibition of nuclear factor (κ)B, p53 activation and ROS stress.3 However, the mechanism of PTL toxicity to T-ALL has not been defined.

Parthenolide has been shown to be very effective against childhood T-ALL in vivo, with elimination of the disease and restoration of murine hemopoiesis in NOD.Cg-Ptkdc<sup>−/−</sup>Il2<sup>−/−</sup>/Sly (NSG) mice.4 However, in mice engrafted with different leukemia initiating cell populations from 2 of 9 T-ALL cases, disease progression was delayed rather than eliminated, indicating variable sensitivity of certain subpopulations to PTL. Reasons for the differences in sensitivity may be due to the effect of the in vivo microenvironment. Bone marrow (BM) stromal cells release cysteine for uptake by chronic lymphocytic leukemia (CLL) cells, driving anti-oxidative glutathione synthesis, which provides protection against ROS generating chemotherapeutic agents, such as fludarabine and oxaliplatin.5

Mesenchymal stem cells (MSC) are key constituents of the BM microenvironment and have been shown to enhance protection against certain drugs in T-ALL cell lines6 and primary samples from patients with B-ALL, acute myeloid leukemia (AML) and CLL.8-10 Co-culture of T-ALL cell lines with MSC enhanced resistance to the anthracycline idarubicin.10 However, the role of ROS in stromal cell mediated protection in childhood ALL has not been reported. As we had previously reported resistance to PTL in T-ALL cases, in this study the cytotoxic and ROS inducing effects of the drug on primary T-ALL cells in the presence of MSC were examined to increase our understanding of PTL resistance.

**Methods**

**T-ALL and normal samples**

Bone marrow samples from 10 children, aged 2-17 years (median 5 years), diagnosed with T-ALL at presentation or relapse were collected with informed consent and approval of University Hospitals Bristol NHS Trust and London Brent Research Ethics Committee (Table 1). Mononuclear cells (MNC) were separated via density gradient centrifugation using Ficoll-Hypaque (Sigma-Aldrich, Gillingham, UK). MNC were suspended in 90% fetal calf serum (FCS, Thermo Scientific, Paisley, UK) and 10% dimethyl sulfoxide (DMSO, Origene Biomedical, Solihull, UK) and stored in liquid nitrogen prior to use. Samples from patients with a range of karyotypic abnormalities, diagnostic age and minimal residual disease (MRD) status were investigated.

Bone marrow from a consenting healthy donor was used as a source of MSC. See the Online Supplementary Appendix for full details of MSC expansion and characterization.

**Cytotoxicity assays**

T-cell acute lymphoblastic leukemia cells were plated in duplicate (for each drug concentration tested) at 1.2x10<sup>5</sup> cells/mL in RPMI 1640 medium (Sigma-Aldrich) containing 20% FCS, 1% L-glut and 1% Pen/Strep, hereafter referred to as suspension medium. Drugs used for assays were: PTL (Enzo Life Sciences, Exeter, UK) at 1-10 μM, N-acetyl cysteine (NAC, Sigma-Aldrich) at 15 mM and 30 μM, and sulfasalazine (SSZ, Sigma-Aldrich) at 300 μM, all prepared in suspension medium. For co-culture experiments, 5x10<sup>4</sup> MSC/mL were seeded per well and left to adhere for 24 hours (h). MSC medium was removed and replaced with suspension media containing 1x10<sup>5</sup> T-ALL cells/mL. T-ALL cells were left to settle for 1 h and then treated with 10 μM PTL with or without 300 μM SSZ, for 24 h. After treatment, non-adherent cells were removed and stained with annexin-V conjugated to fluorescein isothiocyanate (Miltenyi Biotec) for 10 minutes (min). Cells were washed and stained with propidium iodide (PI, Miltenyi Biotec) prior to flow cytometric analysis. See Online Supplementary Figure S1 for details.

For transwell separation experiments, 1x10<sup>5</sup> MSC/mL were seeded per well in MSC media and left to adhere for 24 h. MSC medium was removed and replaced with suspension medium. T-ALL cells were seeded at 2x10<sup>5</sup> cells/mL onto Costar Transwell inserts (0.4 μm pore size, Corning Life Sciences, Ewloe, UK) above the adherent MSC. T-ALL cells were left to settle for 1 h and then treated with PTL with or without SSZ for 24 h. After treatment, cells in transwell inserts were removed and viability was assessed by flow cytometry, as above.

**Reactive oxygen species detection**

Cells were treated with 5 μM of the redox sensitive probe 5-(and-6)-chloromethyl-2,7'-dichlorodihydrofluorescein di-acetate (CM-H<sub>2</sub>DCFDA, Thermo Fisher Scientific) for 30 min at 37°C. Cells were treated with PTL, then immediately analyzed.

---

**Table 1. Patients’ sample characteristics.**

<table>
<thead>
<tr>
<th>Patient N.</th>
<th>Karyotype</th>
<th>Sex</th>
<th>Age (years)</th>
<th>Disease status at biopsy</th>
<th>MRD risk status&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>46XY M</td>
<td>M</td>
<td>7</td>
<td>Diagnosis</td>
<td>Risk</td>
</tr>
<tr>
<td>2</td>
<td>t(1;14) M</td>
<td>M</td>
<td>3</td>
<td>Diagnosis</td>
<td>Risk</td>
</tr>
<tr>
<td>3</td>
<td>46XY M</td>
<td>M</td>
<td>17</td>
<td>Diagnosis</td>
<td>High</td>
</tr>
<tr>
<td>4</td>
<td>t(8;14) M</td>
<td>M</td>
<td>3</td>
<td>Diagnosis</td>
<td>Low</td>
</tr>
<tr>
<td>5</td>
<td>t(11;14), add(17) M</td>
<td>M</td>
<td>2</td>
<td>Relapse</td>
<td>High</td>
</tr>
<tr>
<td>6</td>
<td>RunX1 rearr</td>
<td>M</td>
<td>3</td>
<td>Diagnosis</td>
<td>NA</td>
</tr>
<tr>
<td>7</td>
<td>46XY M</td>
<td>M</td>
<td>6</td>
<td>Diagnosis</td>
<td>Low</td>
</tr>
<tr>
<td>8</td>
<td>add (4), add (9) M</td>
<td>M</td>
<td>14</td>
<td>Diagnosis</td>
<td>High</td>
</tr>
<tr>
<td>9</td>
<td>del (4), del (9) M</td>
<td>M</td>
<td>9</td>
<td>Diagnosis</td>
<td>Low</td>
</tr>
<tr>
<td>10</td>
<td>add (7), add (9), add (14) M</td>
<td>M</td>
<td>4</td>
<td>Diagnosis</td>
<td>Low</td>
</tr>
</tbody>
</table>

*All patients treated on UKALL2011 protocol, except patients 3, 5 and 8 who were treated on UKALL 2003. *Minimal residual disease (MRD) risk status at Day 29, on respective treatment protocols. N: number; M: male.*
over a 90 min period by staining with PI and measuring the median fluorescence intensity (MFI) of the ROS probe by flow cytometry.

**Reduced glutathione detection**

The level of reduced glutathione (rGSH) was detected using the GSH/GSSG-Glo luminescence assay using a plate reader (GloMax, both Promega, Chilworth, UK). Levels of rGSH were quantified in 2x10^4 live T-ALL cells either untreated or treated with 10 μM PTL for 1 h.

**Thiol detection**

The thiol concentration in media was detected using a 5,5′-dithiobis-(2-nitrobenzoic) (DTNB) assay. MSC were plated at 5x10^4 cells/mL per well in MSC medium and allowed to adhere for 24 h. MSC medium was replaced with suspension medium and left for 24 h. Subsequently, 25 μL of MSC conditioned suspension medium was removed and mixed with 100 μL TRIS buffer (0.1M) and 25 μL DTNB (2.5mM, Sigma-Aldrich) solution. The reaction was left for 10 min at room temperature, then the absorbance read at 412nm. The concentration of thiols was quantified using a linear regression equation of the standard curve generated from the absorbance of known standards of cysteine.

**Cysteine transporter, xCT, knockdown in MSC**

Higher thiol levels are a feature of cysteine, which is usually synthesized using a specialized amino acid transport system, known as the xc- system. To investigate whether knockdown of the cysteine transporter, xCT, in MSC affected thiol production small interfering RNAs (siRNAs) for xCT and a non-coding scramble control were transfected into MSC with Lipofectamine 3000 (all Thermo Fisher Scientific) using 20nM siRNA. MSC were plated at 5x10^4 cells/mL per well in MSC media, without Pen/Strep, containing siRNA for 24 h. MSC media was completely removed and replaced with DMEM containing 10% FCS for 24 h. To assess xCT knockdown, mRNA and protein were harvested using a PARIS kit (Thermo Fisher Scientific). (See Online Supplementary Appendix for details.)

**Statistical analysis**

Full details of statistical analyses are provided in the Online Supplementary Appendix.

**Results**

Cytotoxicity is linked to changes in reactive oxygen species (ROS) and reduced glutathione (rGSH) levels can be reversed by N-acetyl cysteine (NAC). (A) T-cell acute lymphoblastic leukemia (T-ALL) samples (patients 1, 3, 5, and 9) were pre-incubated with the ROS sensitive probe CMH2DCFDA and ROS accumulation was measured in live cells over 90 minutes by flow cytometry in response to 10 μM PTL or with PTL+15 mM NAC. Fold change in median fluorescence intensity (MFI) compared to untreated cells is shown; data represent mean ± Standard Deviation (SD). Asterisks represent significant differences between PTL and PTL+15 mM NAC. (B) Dose response curve of patient samples 2, 5, 8, and 10 that had been treated with PTL only (1-10 μM), PTL+30 μM NAC, or PTL+15mM NAC; data represent mean±SD. Top row of asterisks represent significant differences between PTL alone versus PTL+15 mM NAC; bottom row of asterisks represent significant differences between PTL alone versus PTL+30 μM NAC. (C) GSH levels were measured in T-ALL samples (patients 1-9) after treatment with PTL (30 μM) for 1 hour and compared with those in untreated cells. Symbols represent the mean % change from untreated cells in replicate samples. Each symbol represents an individual patient (see Online Supplementary Figure S1 for patient symbol key). Lines represent median and interquartile range. Results were analyzed by two-way ANOVA (A and B) or paired t-test (C). *P≤0.05, **P≤0.01, ***P≤0.001, ****P≤0.0001.
of the antioxidant NAC to PTL treated cells caused a complete block in PTL induced ROS stress over the 90-min period (\( P \leq 0.02 \)) (Figure 1A). NAC (15 mM) reversed PTL induced cytotoxicity, even at the highest dose (10 \( \mu \)M), where T-ALL cells retained a viability of 110±22% compared to 6±3% in cells treated with PTL alone (\( P \leq 0.0001 \)) (Figure 1B). A lower dose of NAC, 30 \( \mu \)M, also significantly reduced PTL cytotoxicity (\( P \leq 0.01 \)), increasing the IC\textsubscript{50} from 2.3 \( \mu \)M to 5.7 \( \mu \)M.

As a further measure of oxidative changes, the levels of the anti-oxidative molecule rGSH were measured in PTL treated cells from samples 1-9. The levels of rGSH detected after 1 h were significantly lower in PTL treated cells with a median of 12% (range 0-58%) compared to untreated cells (\( P = 0.02 \)) (Figure 1C). rGSH levels were higher in the PTL resistant cases (patients 3 and 9).

**MSC protection from PTL and ROS stress**

Mesenchymal stem cells generated from normal BM cells were highly positive for MSC associated markers; CD73 (99±0.3%), CD105 (90±2%), CD90 (99±0.2%) with low expression of hemopoietic cell markers CD45 (0.6±0.2%) and CD34 (0.6±0.5%). MSC protected T-ALL cells from the cytotoxic effects of PTL (78% median survival, range 57-84) in co-culture compared to 8% (range 2-26%) without MSC (\( P = 0.002 \)) (Figure 2A). When T-ALL cells were seeded onto transwell inserts to prevent direct cell contact with MSC, median survival following treatment (58%, range 20-71%) was significantly higher compared to cells treated without MSC support (\( P = 0.05 \)). However, T-ALL survival in transwell cultures was lower than cells in direct contact with MSC (\( P = 0.02 \)) (Figure 2A).

To ascertain whether this MSC conferred resistance to PTL is retained on removal from the supportive environment, T-ALL cells that had been pre-conditioned with MSC for 24 h were treated with PTL without further MSC support. Pre-conditioned cells showed some evidence for enhanced survival compared to cells without conditioning with an increase in IC\textsubscript{50} from 2.6 to 3.3 \( \mu \)M. The increase in PTL resistance in MSC conditioned cells was modest but significant (\( P = 0.04 \)) (Online Supplementary Figure S3).

As ROS levels are associated with PTL cytotoxicity, the ability of MSC to modulate ROS levels was investigated. T-ALL cells conditioned with MSC had significantly lower levels of ROS stress, (ROS MFI 234, range 67-380), compared to those without MSC conditioning (313, range 156-506; \( P = 0.02 \)) (Figure 2B). When T-ALL cells were treated with PTL, ROS levels were significantly increased in both MSC conditioned (\( P = 0.04 \)) and unconditioned (\( P = 0.01 \)) cells. However, the ROS levels following PTL treatment...
were significantly lower in MSC conditioned cells (353, range 200-617) compared to those cultured without MSC (573, range 321-796; *P =0.05) (Figure 2B).

To examine whether these observed protective effects were specific to MSC, T-ALL cells were co-cultured on BM stromal cells and fibroblasts. PTL toxicity was reduced in co-cultures containing stromal cells and fibroblasts but not to the same extent as cultures containing MSC (Online Supplementary Figure S4A). T-ALL cells conditioned with stroma had similar ROS levels to those conditioned with MSC. ROS levels in cells cultured on fibroblasts were similar to those observed in unsupported cultures (Online Supplementary Figure S4B).

Parthenolide is known to lower rGSH levels in leukemia cells, therefore the ability of MSC to modulate this effect was examined. T-ALL cells conditioned with MSC had significantly higher levels of rGSH (median 131nM, range 108-135nM) compared to those without conditioning (82nM, range 60-99nM; *P=0.03). Following PTL treatment, rGSH levels were significantly decreased in both MSC conditioned (*P=0.04) and unconditioned (P=0.0003) cells, indicating reduced anti–oxidant activity. However, rGSH levels were still higher in MSC conditioned cells (median 87nM, range 66-117nM) compared to those without MSC conditioning (26nM, range 7-35nM; **P=0.01) (Figure 2C).

As cysteine is crucial for GSH synthesis in leukemia, it is possible that MSC release cysteine into the surrounding media, thereby moderating the anti-oxidant effects of PTL. A thiol detection assay was used, since cysteine contains a free thiol group, to investigate cysteine release. Thiol concentrations were significantly higher in media removed from MSC cultures (91±17 μM) compared to background media levels (62±17 μM; *P=0.0002), representing a 1.5-fold increase over background (Figure 2D).

Blocking thiol release overcomes protective effect

The thiol functional group plays a major role in intracellular anti-oxidant defenses. Cysteine residues eliminate ROS, usually by converting them to 2H2O or H2O and O2, and reduce oxidized protein thiols. SSZ, an inhibitor of the cystine uptake antipporter xc−, was used to inhibit cysteine uptake and block subsequent cysteine generation and release in MSC, to determine what effect this would have on the protection conferred by MSC. The combination of PTL and SSZ significantly reduced the median viability of T-ALL cells with MSC conditioning to 38% (range 11-42%) compared to PTL (71%, range 59-95%; **P=0.002) or SSZ alone (88%, range 69-109%; *P=0.04) (Figure 3A). The treatment combination reduced viabilities to levels comparable to those observed in cells treated with both agents without MSC conditioning (16%, range 5-24%). In the absence of MSC, SSZ had low cytotoxicity (92% median survival, range 77-102%) and when combined with PTL there was no significant difference in toxicity compared to PTL alone (P=0.9). Addition of PTL+ SSZ to T-ALL cells conditioned with BM stroma or fibroblasts also reduced leukemia cell survival, although to a lesser extent (Online Supplementary Figure S4C).

Interestingly, the effects of combining these agents were more remarkable in samples that were relatively resistant to PTL. Fifty-six percent of cells from patient 9 survived treatment with PTL, but viability was reduced to only 6% when PTL was combined with SSZ. Leukemia burden in NSG mice engrafted with this sample was significantly reduced when treated with PTL and SSZ compared to placebos.
Using SSZ alone and in combination with PTL had minimal effects on normal hemopoietic cells (Online Supplementary Figure S6).

In transwell experiments, using SSZ with PTL abrogated the protective effects of MSC support, reducing T-ALL survival from 58% (range 20-71%) to 14% (range 9-43%; \( P=0.01 \)) (Figure 3B). Media harvested from cultures of SSZ treated MSC contained thiol levels similar to background media (53±19 \( \mu \)M vs 61±10 \( \mu \)M, respectively). In contrast, media from untreated MSC contained significantly higher thiol levels (90±11 \( \mu \)M; \( P=0.005 \)) compared to media from SSZ treated cells, representing a 1.5-fold increase over background media (Figure 3C).

As a second approach to block the cystine uptake antiporter xc\(^-\), and thereby prevent cysteine release from MSC, xc\(^-\) was targeted with siRNAs. Relative gene expression levels of xc\(^-\) were significantly lower in MSC 48 h after transfection with either siRNA 1 (17±9%) or siRNA 2 (9±4%) compared to scramble control siRNA (\( P\leq0.0001 \)) (Online Supplementary Figure S7A). Protein available from xc\(^-\) siRNA-1 showed a significant reduction in xc\(^-\) expression (3±0.1%) relative to scramble control levels quantified by western blotting densitometry (\( P=0.002 \)) (Online Supplementary Figure S7B).

The viability of PTL treated T-ALL cells, co-cultured with MSC treated with xc\(^-\) siRNA-1 (27%, range 3-28%) or -2 (35%, range 24-42%), was significantly lower than cells co-cultured with scramble control MSC (49%, range 46-72%; \( P<0.05 \)) (Figure 4A). As siRNA-1 provided the best reduction in MSC protection, it was selected for further experiments. When transwell inserts were used to separate xc\(^-\) knockout MSC from T-ALL cells, the median viability after PTL treatment was significantly lower (15%, range 7-30%) compared to scramble control MSC cells (24%, range 15-74%; \( P=0.05 \)) (Figure 4B). Scramble control MSC media had a significantly higher thiol concentration (86±2 \( \mu \)M) compared to media harvested from xc\(^-\) knockout MSC (46±1 \( \mu \)M; \( P=0.002 \)) (Figure 4C). Media from xc\(^-\) knockout MSC did not show any increase in thiol levels above background media (53±1 \( \mu \)M).

To confirm that the observed effects of interfering with xc\(^-\) were a result of blocking cystine uptake, and not a result of toxicity, the viability and morphology of treated MSC were assessed. Treatment with PTL, SSZ or both agents in combination had no effect on the viability and confluency of MSC (\( P\geq0.37 \)) (Figure 5A) nor did siRNA treatment (\( P\geq0.82 \)) (Figure 5B and Online Supplementary Figure S8). This was also confirmed by flow cytometry (Figure 5C).

**Discussion**

To date, PTL is the only drug that has been shown to be capable of completely eradicating childhood ALL in NSG xenografts, as a single agent.\(^8\) Most studies, using such models, report reduction in leukemia burden but levels often increase on cessation of treatment. Consequently, there is much interest in the application of PTL for cancer therapy. Several groups are developing strategies to improve the bioavailability of PTL, without having detrimental effects on its pharmacokinetic properties.\(^16\)\(^-\)\(^19\) PTL can be successfully sequestered into nanoscopic vectors which can achieve equivalent toxicity to unmodified PTL.\(^16\)\(^-\)\(^19\) Nanoparticle PTL formulations can be used *in vivo* at 40-fold lower doses with 20-fold lower administration fre-
frequency than standard PTL. Some of these delivery systems are inexpensive and formulations are readily scalable. Consequently, they should facilitate the use of PTL at clinically relevant doses. However, the efficacy of many therapeutic drugs may be compromised by the host BM microenvironment. Therapy-induced niches, which protect leukemia cells against standard first-line induction agents, have been described in ALL. In the present study, we investigated whether resistance to PTL, reported in a minority of T-ALL cases, may be conferred by BM-derived MSC.

In vitro PTL treatment reduced viability of T-ALL cells to less than 50%, confirming previous results in a separate cohort of pediatric cases. While there were differences in the responses of individual patient samples, there was no correlation between PTL cytotoxicity with karyotype or MRD risk status, which may be a result of the heterogeneous nature of this disease. PTL was shown to increase ROS stress and lower rGSH levels in T-ALL. The anti-oxidative compound NAC blocked ROS upregulation and ROS stress and lower rGSH levels, suggesting that MSC increased the anti-oxidative stress capacity of T-ALL cells and thereby conferred resistance to PTL. To eliminate variability, MSC were generated from a single normal BM donor, allowing a direct comparison of the effects on the patients’ samples. This may have introduced bias but our results are consistent with those reported in CLL. Chemo-protection was derived from the generation and release of cysteine by stromal cells for uptake by CLL cells. The BM microenvironment can also play a role in leukemia drug resistance, mediated by a diverse set of mechanisms, both by direct cell adhesion and/or soluble factor release.

The protective effect provided by NAC may be attributed to the role of cysteine as the rate limiting amino acid in rGSH production. Reduction of cysteine to cysteine and subsequent supply to leukemia cells is crucial for GSH synthesis in these cells. GSH levels elevate the anti-oxidative capacity of cells, which may provide protection against PTL. In AML and breast cancer, populations of cells endure lower levels of ROS stress and these cells are more resistant to therapy. It is unclear whether ROS is actively eliminated by rGSH or if PTL directly interacts with rGSH, leading to a reduction in anti-oxidative capacity and increased ROS accumulation. One function of rGSH is to detoxify cells from reactive electrophiles, which PTL contains in the form of an α,β-unsaturated carbonyl group. PTL can directly interfere with rGSH synthesis in AML, causing depletion of rGSH, allowing ROS levels to increase. Whether PTL directly blocked rGSH synthesis was beyond the scope of this study. However, it is evident that ROS levels increased and rGSH levels dropped, putting the cells under higher levels of stress, which is likely to drive apoptosis.

As these data indicate that ROS levels may be linked to PTL cytotoxicity, it is possible that the BM microenvironment provides resistance to drug activity by protecting leukemia cells from ROS stress. CIL cells had decreased sensitivity to the ROS inducing agents fludarabine and oxiplatin when co-cultured with BM stromal cells. Chemoprotection was derived from the generation and release of cysteine by stromal cells for uptake by CIL cells. The BM microenvironment can also play a role in leukemia drug resistance, mediated by a diverse set of mechanisms, both by direct cell adhesion and/or soluble factor release.

In this study, the protective effects of MSC against PTL toxicity were related to decreases in ROS stress and preserved rGSH levels, suggesting that MSC increased the anti-oxidative stress capacity of T-ALL cells and thereby conferred resistance to PTL. To eliminate variability, MSC were generated from a single normal BM donor, allowing direct comparison of the effects on the patients’ samples. This may have introduced bias but our results are consistent with those reported in CLL. These data also suggest that protection is provided by one or more secretable factors, as preventing direct contact with MSC still provided T-ALL cells with protection against PTL. However, protection was not to the same extent as observed in direct contact, suggesting involvement of other factors.

Mesenchymal stem cell conditioned media contained significantly higher levels of thiols, an important feature of the anti-oxidative compound cysteine, compared to unconditioned media. A key mechanism by which normal cells synthesize cysteine is via intake of extracellular cysteine.
using a specialized amino acid transport mechanism known as the xc\(^{-}\) system.\(^{14}\) The system specifically mediates cysteine and glutamate exchange across the cell membrane and is comprised of two subunits: the light chain transporter subunit SLC7A11 (or xCT) and the heavy subunit SLC3A2 (4F2hc).\(^{30,31}\) Cysteine is unstable outside of cells and is rapidly oxidized to form a disulphide bridge with another cysteine amino acid, forming cystine. However, upon entry into the cell, cystine enters a stronger reducing environment and is converted into cysteine. Blocking xc\(^{-}\) activity with a small molecule inhibitor, SSZ,\(^{32}\) or by knockdown of the xc\(^{-}\) light chain component xCT, prevented thiol release and significantly reduced MSC protection to levels close to those observed in cells without MSC support. In addition, leukemia cell survival in a PTL resistant case was significantly decreased when SSZ or xCT siRNA were used to block xc\(^{-}\) activity. Blocking PTL and SSZ treatment in vivo resulted in significantly reduced leukemia burden in engrafted NSG mice compared to controls. Similar results have been reported in BCP-ALL, where cell viability was diminished in a subset of patients following treatment with cysteine dioxgenase, which catalyzes conversion of cysteine into cystine sulfonic acid, thereby bypassing GSH synthesis.\(^{35}\) Together, these findings indicate that MSC release cysteine, which confers a survival advantage on leukemia cells. The functional contribution of thiols was further confirmed by the demonstration that T-ALL cells exposed to the thiol containing compound NAC, at an equivalent thiol concentration to that in MSC conditioned media, were more resistant to PTL. The fact that there was still a small protective effect against PTL after interfering with the xc\(^{-}\) system, indicates that cysteine release may not be the sole protective mechanism in these experiments. A recent study found that ALL cells release extracellular vesicles that can be taken up by MSC, causing a shift to glycolysis metabolism. The switch to glycolysis led to an increased release of the metabolite lactate, which might be used as an additional source of energy by leukemia cells and confer chemoresistance.\(^{34}\)

Figure 6. Proposed mechanism of action of parthenolide (PTL) and the protective effect provided by mesenchymal stem cells (MSC). PTL causes apoptosis by increasing reactive oxygen species (ROS) stress and decreasing reduced glutathione (rGSH) resulting in death of T-cell acute lymphoblastic leukemia (T-ALL) cells (A). MSC express high levels of the cystine glutamate antiporter xc\(^{-}\), facilitating entry of the antiporter protein xCT. Extracellular cystine is taken up by MSC and reduced into cysteine. Cysteine is released into the extracellular space for uptake by T-ALL cells, blocking ROS induction and preventing apoptosis (B). Blocking the xCT system using sulfasalazine (SSZ) or siRNA reduces protective effects provided by MSC (C).

While MSC are a fundamental component of the BM niche, and contribute to its formation in vivo, other stromal cells may have a role in leukemia maintenance and protection from therapeutic agents. We demonstrated that BM stromal cells and fibroblasts also confered protection against PTL. ALL cells themselves can alter the endosteal and vascular compartments of the niche\(^{12}\) and induce apoptosis of osteoblast cells.\(^{37}\) The remodelled niche is dynamically transient and on exposure to chemotherapeutic agents, ALL cells can release CCL3 and cytokines TGF-\(\beta\)1 and GDF15, conferring chemoresistance by activating the TGF-\(\beta\) signaling pathway.\(^{37}\) It may be possible to overcome this resistance by disrupting the interactions between leukemia cells and the BM environment.

This is the first report demonstrating MSC provide a protective effect to T-ALL cells against PTL. Moreover, we have shown that targeting the xc\(^{-}\) system can overcome this effect (see overview in Figure 6), adding to the evidence that targeting xc\(^{-}\) enhances the efficacy of anti-cancer agents in leukemias\(^{9}\) and in several solid cancer models.\(^{38-41}\) MSC viability was unaffected by PTL and targeting xc\(^{-}\) resulted in damage to this important element of the BM environment\(^{42-45}\) should be minimal. A logical progression of this work will be determining whether the effects of PTL, or indeed current chemotherapeutic agents, in vivo can be enhanced using an xc\(^{-}\) inhibitor like SSZ. Furthermore, with developments in nanoscopic drug delivery vectors, it may be possible to use dual loaded (PTL+SSZ) nanoveccors to overcome selective patient resistance towards one drug and the chemoprotective effects of the leukemia microenvironment in vivo.

Acknowledgments
The authors would like to thank Dr Jeremy Hancock, Mr Paul Virgo and staff of Bristol Genetics Laboratory, Southmead Hospital for excellent technical assistance. We also thank Elinor Curnow, Statistics and Clinical Studies, NHS Blood and Transplant, consultants and oncology staff at Bristol Royal Hospital for Children. We are grateful to the patients and their families who gave permission for their cells to be used for research. This work was supported by a generous donation from Mr Richard Cunningham and by grants from the Department of Health and NHS Blood and Transplant.
References