Metabolic plasticity in blast crisis-chronic myeloid leukaemia cells under hypoxia reduces the cytotoxic potency of drugs targeting mitochondria

Luciana S. Salaverry · Tomás Lombardo · María C. Cabral-Lorenzo · Martin L. Gil-Folgar · Estela B. Rey-Roldán · Laura I. Kornblihtt · Guillermo A. Blanco

Received: 7 April 2022 / Accepted: 16 June 2022
© The Author(s) 2022

Abstract

Metabolic reprogramming (MR) influences progression of chronic myeloid leukaemia (CML) to blast crisis (BC), but metabolic programs may change transiently in a second dimension (metabolic plasticity, MP), driven by environments as hypoxia, affecting cytotoxic potency (CPot) of drugs targeting mitochondria or mitochondria-related cell stress responses (MRCSR) such as mitophagy and mitochondrial biogenesis. We assessed mitochondrial membrane potential (MMP), mitochondrial mass (MM), apoptosis, glucose uptake (GU), and CPot of arsenic trioxide (ATO), CCCP, valproic acid (VPA), vincristine (VCR), Mdivi1, and dichloroacetic acid (DCA) in CML BC cells K562 (BC-K562) under hypoxia through flow cytometry, and gene expression from GEO database. About 60% of untreated cells were killed after 72 h under hypoxia, but paradoxically, all drugs but ATO rescued cells and increased survival rates to almost 90%. Blocking mitophagy either with VCR or Mdivi1, or increasing mitochondrial biogenesis with VPA enhanced cell-survival with increased MM. DCA increased MM and rescued cells in spite of its role in activating pyruvate dehydrogenase and Krebs cycle. Cells rescued by DCA, VPA and CCCP showed decreased GU. ATO showed equal CPot in hypoxia and normoxia. MP was evidenced by differential expression of genes (DEG) under hypoxia related to Krebs cycle, lipid synthesis, cholesterol homeostasis, mitophagy, and mitochondrial biogenesis (GSE144527). A 25-gene MP-signature of BC-K562 cells under hypoxia identified BC cases among 113 transcriptomes from CML patients (GSE4170). We concluded that hypoxic environment drove a MP change evidenced by DEG that was reflected in a paradoxical pro-survival, instead of cytotoxic, effect of drugs targeting mitochondria and MRCSR.

Keywords Metabolic reprogramming · Arsenic Trioxide · Dichloroacetate · Gene expression profiling · Valproic acid · Glucose uptake

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s12672-022-00524-y.
1 Introduction

Chronic myeloid leukaemia (CML) is caused by a chromosomal translocation that results in the expression of the fusion protein bcrabl1 with kinase activity [1]. The course of CML was radically modified with the introduction of tyrosine kinase inhibitors (TKIs) such as imatinib. However, imatinib does not cure the disease and interruption of treatment usually leads to relapse. In some cases, CML evolves from chronic phase (CP) into more advanced forms, such as accelerated phase (AP) and blast crisis (BC) [2], with resistance to TKIs due to mutations in the bcrabl1 protein. However, in many cases this type of mutation is not detected and the cause of progression remains elusive.

Metabolic reprogramming (MR) is considered a neoplastic hallmark, and involves the expression of genes associated with aberrant metabolic programs compared to the cells of origin [3]. MR is a phenotypic trait or dimension that evolves along disease progression. For example, metabolic pathway genes may show different profiles of expression in BC cells, AP cells, and CP cells [4]. However, alternative metabolic programs are not permanently active but may change in a second phenotypic dimension according to the microenvironment. This rather transient changes are often referred to as metabolic plasticity (MP), and allow neoplastic cells to tolerate temporary harsh conditions such as lack of oxygen or nutrients (glucose, or amino acids such as glutamine) [5]. For example, mitochondrial and metabolic pathway genes may show different profile of expression in BC cells under hypoxia compared to the same cells under normoxia. Therefore, MR and MP are independent phenotypic dimensions, with the latter being driven by transient changes in the microenvironment. Increasing MP is supported by anabolic and catabolic pathways that largely interconnect at the mitochondrial level. The most direct example of MP is the shift to glycolysis as the main source of ATP under hypoxia, to compensate reduced oxidative phosphorylation (OXPHOS) at the mitochondria. However, mitochondria can assist biosynthetic pathways, where the Krebs cycle functions to provide metabolites that are exported to the cytoplasm for the synthesis of nucleotides, fatty acids, cholesterol, amino acids, and antioxidant molecules [3, 6].

Mitochondria initiate apoptosis and are the target of most cytotoxic drugs [7]. Being at the intersection of metabolic and apoptosis control, mitochondria may influence the threshold of cytotoxic drugs, depending on the microenvironment and the metabolic program adopted by cells [8]. Cytotoxic drugs must alter mitochondrial function sufficiently enough to reach the threshold of apoptosis initiation and kill cells. For this reason, metabolic reprogramming, and more specifically the metabolic pathways regulated by mitochondria, can significantly modify the effect of cytotoxic drugs [9]. In other words, MR along disease progression, and MP between different environments, can not only be observed at transcriptomic, proteomic or metabolic dimensions, but can also be exposed by altered drug potency in harsh environments such as hypoxia or lack of nutrients.

The physiological response to hypoxia in normal cells is mediated by hypoxia-inducing-factor (HIF). HIF-targeted genes are known to increase glycolysis as a source of ATP, and reduce mitochondrial mass (MM) through increased mitophagy [10]. The classical view considers that mitochondria cannot produce ATP under hypoxia, and therefore their rapid elimination through mitophagy is necessary to prevent their failure, since a failing OXPHOS may lead to mitochondrial apoptosis [11]. It is often assumed that mitochondrial failure due to lack of O₂ could increase sensitivity to drugs that activate mitochondrial apoptosis. This has established a general principle for blocking the adaptation to hypoxia of neoplasms, which consists of actions such as inhibiting mitophagy, inducing an increase in MM by targeting mitochondrial biogenesis, or forcing the resumption of OXPHOS with metabolic drugs, therefore leading to massive mitochondrial failure and onset of apoptosis [12, 13]. However, the metabolic change induced by hypoxia in advanced leukaemia could have an influence on mitochondrial apoptosis. One way of exploring the impact of microenvironment-induced MP is to explore the cytotoxic potency of drugs under hypoxia. Hypoxia below 1% induces a metabolic change, since mitochondria cannot produce enough ATP and must resort to glycolysis. CCCP is a compound that blocks OXPHOS by uncoupling electron transport chain (ETC) causing significant mitochondrial damage and inducing apoptosis [14, 15]. In the presence of CCCP it is not possible to obtain ATP via OXPHOS and the only alternative to produce ATP is glycolysis. Arsenic trioxide (ATO) is a widely studied compound that targets mitochondria and also inhibits OXPHOS, forcing glycolysis as an alternative source of ATP [16]. However, it does so not only at the level of ETC, but also by inhibiting key enzymes of the Krebs cycle such as pyruvate dehydrogenase (PDH) and 2-oxoglutarate dehydrogenase (2OGDH) [17]. Vincristine (VCR) and Mdivi1 are two drugs that inhibit mitophagy, but differ in the mode of inhibition. Mdivi1 blocks mitochondrial fission, which in turn is the first step of mitophagy, and consists of separating those mitochondria of low quality or with bioenergetic failure [18, 19]. VCR inhibits the polymerization of microtubules, preventing the transport of damaged mitochondria into lysosomes, and
in particular the fusion of “mitophagosomes” with lysosomes, which is the final step of mitophagy [12, 20]. Valproic acid (VPA) is a histone deacetylase inhibitor that activates the expression of PGC1α, the main transcription factor of mitochondrial biogenesis [21], while dichloroacetate acid (DCA) is an inhibitor of pyruvate kinase-1 (PDK1) and offsets HIF-driven response to hypoxia by restoring the function of PDH, the Krebs cycle and OXPHOS [22]. All these drugs should be very toxic for adaptation to hypoxia considering the classical physiological principles of survival under low oxygen tension.

The objective of this work was to expose BC-K562 cells to conditions of hypoxia (< 1% O₂) in order to force a metabolic change, and explore the cytotoxic effect of the six above-mentioned drugs that target mitochondria, or mitochondria-related cell stress responses (MRCSR) such as mitochondrial biogenesis, mitophagy, and cell metabolism (ATO, CCCP, Mdivi1, VCR, VPA and DCA). Our results showed several paradoxical effects of these drugs, contrasting the classical view. We interpret that these effects are caused by both the MR of BC cells, and MP driven by the hypoxic environment. We support our interpretation exploring differential expression of genes (DEG) related to mitochondria, MRCSR and metabolism in BC-K562 cells under hypoxia, as an indication of MP (MP-signature). In addition, we investigated if these genes were also differentially expressed in BC cells from CML patients compared to CP and AP patients as an indication of overall MR during CML progression (MR-signature). From the analysis of two public available gene expression data sets, we concluded that MP of BC-K562 under hypoxia promoted mitochondrial functions not related to OXPHOS but to mitochondrial anabolic pathways and MRCSR, and that these changes could be the cause of the paradoxical effect of drugs on BC-K562 under hypoxia.

2 Materials and methods

2.1 Cell model, culture conditions and reagents

The study was conducted on the cell line K562 that was originally derived from a sample of the pleural effusion of a 53-old female CML patient in BC eight days prior to her death in 1970 [23]. This cell line is positive for the BCR:ABL fusion gene (Philadelphia chromosome) and is included in a panel of the 60 cell lines selected by the National Cancer Institute of USA to identify and characterize novel anti-cancer compounds [24]. K562 cells were purchased from ATCC and grown in RPMI 1640 with 10% foetal bovine serum (FBS) at 37 °C in a 5% CO₂ atmosphere. Cells were subcultured at 1:2 ratios every 2 to 3 days with a viability greater than 95%. For the experimental assessment of drugs, K562 cells were decanted in 24-well culture plates at 0.5 × 10⁶/ml. Serial dilutions of the corresponding single drug or combinations were added in triplicate. RPMI was added as untreated control. The plates were incubated at 37 °C with 5% CO₂ at ambient air (21% O₂) and were denoted as “normoxia” experimental conditions.

A second series of experiments that we denoted as “hypoxia” experimental conditions was conducted in similar culture plates but within a Modular Incubator Chamber (MIC-101) from Billups Rothenberg, Inc. (Del Mar, USA) as described previously [25]. The chamber was attached to a flow meter from the same manufacturer (SFM3001; 3–25 LPM adjustable flow rate) to regulate the gas flow. To attain hypoxia conditions after adding the drugs, the cells were immediately placed in the chamber and flushed for 15 min with a gas mixture of 5% CO₂ and 95% N₂. The chamber was sealed and placed at 37 °C in a conventional cell incubator, ensuring the O₂ concentration is maintained below 1.0% up to 72 h [26, 27].

The classical view of survival under hypoxia considers that mitochondria is superfluous as source of energy under low O₂ tension, and may pose severe risks of cell death due ETC failure. Assuming that blocking elimination of mitochondria or even increasing mitochondrial network would be severely deleterious for cell survival under hypoxia, we exposed BC-K562 cells to conditions of hypoxia in order to force a metabolic change, and explore the cytotoxic effect of six drugs that target mitochondria and MRCSR, such as mitochondrial biogenesis (VPA), mitophagy through microtubule transport inhibition (VCR) and blockage of mitochondrial fission (Mdivi1), disruption of ETC and OXPHOS without compromise of Krebs cycle (CCCP), disruption of ETC and Krebs cycle (ATO), and disruption of Krebs cycle with offset of HIF1-mediated response by increased PDH activity (DCA).

Carbonyl cyanide 3-chlorophenylhydrazone (CCCP), and 3-(2,4-Dichloro-5-methoxyphenyl)-2,3-dihydro-2-thioxo-4(1H)-quinazolinone (Mdivi-1) were from Sigma (Buenos Aires, Argentina). Stock solutions were prepared in dimethyl-sulfoxide (DMSO) at 80 mM and 5 mM concentration respectively. Valproic acid was from Casasco Pharmaceuticals (Argentina) and was prepared freshly as a 100 mM stock solution in RPMI-1640. Vincristine was kindly provided by Laboratorio Filaxis (Argentina). Arsenic trioxide (ATO) was a kind gift from Varifarma (Argentina). Fresh stock solutions of 1 mM ATO were prepared before every assay. DCA was from Sigma-Aldrich (Buenos Aires, Argentina). Fresh stock solutions of
1M DCA in RPMI-1640 were prepared before every assay. RPMI-1640, penicillin, streptomycin, tetramethylrhodamine-ethyl-ester (TMRE), nonyl-acridine orange (NAO), and 2-deoxyglucose analog, 2-[N-(7-nitrobenz-2-oxa-1,3-diaxol-4-yl) amino]-2-deoxyglucose (2-NBDG) were purchased from ThermoFisher, and prepared as a 10mM stock solution in DMSO.

2.2 Labelling of cells with fluorescent probes

We employed a flow cytometry multiparametric method to simultaneously assess mitochondrial mass (MM), mitochondrial membrane potential (MMP), and cell death in single cells as described previously [15]. Briefly, the MMP was evaluated by the potentiometric probe TMRE [12]. Cells were incubated in 0.05μM TMRE for 20 min at 37°C. Damage to cell membrane was detected by labelling with 1μM propidium iodide (PI). The MM was evaluated with nonyl-acridine orange (NAO) [28]. NAO was prepared as a 5mM stock solution in ethanol, and diluted to 100nM in PBS to stain cells during 15min at 37°C. The cells were washed once in PBS and analysed by flow cytometry. To determine glucose uptake cells were labelled with 25μM 2-NBDG during 40min and washed once in PBS before flow cytometry analysis.

2.3 Flow cytometry analysis

All samples were evaluated in a Partec PAS III flow cytometer equipped with a 20mW 488nm argon blue laser (Partec, GmbH, Münster, Germany). A total of 20,000 cells were analysed in each sample. Fluorescence from NAO and 2-NBDG was collected through a 520/10nm bandpass (BP) filter (FL1). Fluorescence from TMRE was collected with a 590/20nm BP filter (FL2) and PI was collected with 670/20nm BP filter. Flow cytometry data analysis was performed with Flomax software (Partec, Germany) and Flowing Software 2.5.1 (Turku Centre for Biotechnology, Finland). Unsupervised classification of labelled cells through flow cytometry self-organized maps (FlowSOM) algorithm [29], and visualization of cells in a five-dimensional space (FSC, SSC, FL1, FL2 and FL3) through t-distributed stochastic neighbour embedding (tSNE) plots [30], was conducted with FlowJo software 10.5.0 (BD, USA) using the standard plugins.

2.4 Assessment of the mean cytotoxic effect of single drugs

We used a quantal model of dose–response function, where the number of dead and live cells scored by flow cytometry was used to define two ratio measures. $Fa$ was the fraction of cells affected by cytotoxic drugs, and $Fu$ was the fraction of live cells, with $Fu = (1−Fa)$. By linear regression, we obtained the slope and the intercept of the equation $\log(Fa/Fu) = m. \log(D)−m. \log(Dm)$, where D was the dose of the drug tested. With the slope m and the median effect dose Dm we obtained the median effect equation for each individual drug as: $D = Dm (Fa/(1-Fa))^{1/m}$. With this formula, we estimated the dose D of drug causing the death of fraction $Fa$ of cells after 72h. Therefore, we referred to the $Fa$ values as the cytotoxic effect (EC) and we denoted EC50 as the dose that causes death in 50% of cells [25, 31].

2.5 Differential gene expression

To explore changes in the expression of genes related to mitochondria and metabolism that could be induced by hypoxia in BC-K562 cells, we compared the transcriptomes of BC-K562 cells under normoxia and hypoxia, using raw data from an RNAseq dataset that became available recently in GEO database (GSE144527). This dataset includes six transcriptomes obtained from BC-K562 cells exposed to 1% O2 and 21% O2 during 72h [32].

After obtaining a list of 25 genes related to mitochondria and metabolism from our analysis on the GSE144527, whose expression characteristically changed under hypoxia (MP-signature), we further explore if these genes were also differentially expressed in BC cells from CML patients. For this purpose we used another transcriptomic dataset from a large cohort of CML patients (GSE4170), also deposited in the GEO database. This dataset includes 113 transcriptomes from CML patients corresponding to 62 cases of CP, 17 cases of AP and 33 cases of BC [33]. We used the conventional bioinformatics tools GEOquery, limma (Linear Models for Microarray and RNA-Seq Data) and edgeR from R bioconductor to evaluate DEG and detect functional groups of genes associated with metabolism and mitochondrial function. We listed the top-ranked individual DEG using lmFit, eBayes and TopTable functions of the R limma package. Next, we listed the top-ranked Gene Ontology (GO) sets with DEG in terms of cellular component (CC), biological processes (BP), and molecular functions (MF) using GOANA functions of the limma R package, and we listed the top-ranked pathway gene sets with DEG from the Kegg pathway database using the KEGGA function of the limma R package. The entire fitted linear
model (fit object) as obtained with lmFit and eBayes functions of limma package, containing all genes probed, was used as input for GOANA and KEGGA functions, using the default FDR cut-off parameter value of 0.05 [34].

2.6 Gene set enrichment analysis and hierarchical clustering

We explored coordinated expression of genes known to be involved in particular functions through the ranks-based methods provided by the Gene Set Enrichment Analysis (GSEA) software platform, which includes a Molecular Signature Database (MSigDB) with eight major collections of gene sets. We used the Hallmark collection of gene sets that consists of 50 sets of well-defined biological states or processes with little overlap between them, to obtain an unbiased overview of cell states under hypoxia and normoxia in BC-K562 cells.

The GSEA analysis is not restricted to the list of genes with differential expression above a certain level of statistical significance, but instead performs an evaluation using all genes for which the dataset has available RNA expression values. In this way GSEA can track even several minor changes occurring simultaneously in specific functional gene lists, using a statistical method based on ranks. The ranked differential expression list was built using the default ranking metric “signal-to-noise ratio” of the GSEA software, which is defined as S2N = (x_1 − x_2) / (s_1 − s_2), where for each gene, x_1 and s_1 corresponded to the average and standard deviation of gene expression of three samples under hypoxia, and x_2 and s_2 corresponded to the average and standard deviation of three samples under normoxia [35]. Nominal p-values and q-false discovery rates (FDR) probability measures were used as statistical significance values of gene set enrichment, as calculated by GSEA software [35]. We restricted the analysis to 4362 genes (from the large set of 21,950 genes) whose expression was consistently detected in all 6 samples of GSE144527 (three replicates of BC-K562 cells in normoxia, and three replicates in hypoxia), considering genes with at least 10 counts from the RNAseq assessment in all 6 samples. In this way, we constrained the exploration of enrichment to genes with clear expression in BC-K562 cells under normoxia and hypoxia.

We further defined gene sets potentially associated to metabolic plasticity from GSEA and DEG analysis, and we conducted unsupervised classification through hierarchical clustering. Classification was conducted on gene expression values in transcriptomes from GSE144527 and GSE4170 using hclust and dendextend packages from R-Bioconductor, for computing the Euclidean distance and applying the complete linkage method. Additional R packages used for hierarchical clustering analysis and heatmaps production were gplots, factoextra, and dendextend [36].

2.7 Statistics

In DEG analysis a p-value below 0.05% was considered significant. An adjusted p-value < 0.05 was computed through the Benjamini-Horchberg method to consider false positives rates (FDR) due to multiple comparisons. Bars in bar graphs in figures and supplemental materials indicate SE of the mean fluorescence intensity or fraction of cells from three replicates. In all cases, bar graphs show one experiment representative of at least three. Spearman-rank correlation analysis and contingency tables analysis were performed with GraphPad Prism 7.0 (GraphPad Software, Inc.).

3 Results

3.1 Cytotoxicity of ATO on BC-K562 cells under normoxia

We evaluated drug-induced changes in MM, MMP, and apoptosis on BC-K562 cells using a flow cytometry triple-staining method with NAO-TMRE and PI. NAO produces bright fluorescence when bound to reduced-cardiolipin (rdCL) at the mitochondria, but emits low fluorescence when bound to oxidized-cardiolipin (oxCL) [28]. The abrupt drop in NAO intensity with collapsed MMP (TMRE low) is an indicator of ongoing apoptosis with mitochondrial outer membrane permeabilization (MOMP), massive oxidation and passage from rdCL to oxCL and release of cytochrome c. Increasing doses of ATO at 72 h caused a gradual decline in MMP with rdCL, followed by an increase in oxCL without cell membrane damage, and finally loss of viability becoming PI-positive. These transitional dose-dependent changes can be observed in bivariate plots as shown in Fig. 1a.

Considering the fraction of dead and live-apoptotic cells with oxCL (Q1 + Q3 in Fig. 1a), we quantified the cytotoxic potency using the median effect equation to compute the EC50 (Fig. 1b). We observed a similar dose-dependent reduction in MMP in 72 h samples (Fig. 1c). We observed increased MM (mean fluorescence intensity in Q2 and Q4) up to 10 μM...
yet in both situations the viability rose to about 90% (Fig. 3b). Therefore, MMP was needless for survival under hypoxia. Conversely, CCCP reduced glucose uptake within the dose range where cell survival was increased (4-10 μM). Detected at 3 μM CCCP or below, almost disappearing at higher CCCP doses (Supp. Fig. S1).

At 5 μM CCCP, rescued cells were all in the low MMP group (matching Q4). In both cases rescued cells showed high NAO fluorescent signal (rdCL content). Early alive apoptotic cells matched the Q3 population in quadrant analysis, and were even detected at 3 μM CCCP or below, almost disappearing at higher CCCP doses.

A large fraction of viable cells in samples treated with 5 μM CCCP and above, still showed high NAO fluorescence signal (rdCL) with collapsed-MMP (Q4 in Fig. 2b) after 72 h treatment. These cells were able to survive in spite of the restriction posed to OXPHOS by collapsed MMP and uncoupled ETC caused by CCCP. Therefore, cells survived with collapsed MMP and rdCL content, and even restored the MMP 72 h later (progressing from Q4 back to Q2, Fig. 2a, b).

An increased MM was observed at very low dose ranges (< 3 μM CCCP), while at higher dose ranges MM was reduced (Fig. 2d). Massive collapse of MMP restrains ETC and OXPHOS, and should force the activation of glycolysis as a compensatory source of ATP. However, we observed a progressive reduction in glucose uptake over the range 0-10 μM CCCP (Fig. 2e, f).

**3.3 CCCP rescued BC-K562 cells from hypoxia-induced death**

Exposure to hypoxia for 72 h induced cell death in more than 50% of cells considering membrane damage and oxCL, and live cells in early apoptosis with oxCL (Q1 + Q3) (Fig. 3a, b). These figures were coincident to a previous study in BC-K562 cells under 1% O2, even though the authors used the method of AnnexinV with PI to quantify apoptosis [37]. Paradoxically, the mitochondrial damage induced by CCCP was beneficial for cell survival under hypoxia in a dose-dependent manner in the 0–10 μM range (Fig. 3a, b). For example, with 10 μM CCCP, there were 42% of viable cells with high MMP, and 43% of viable cells with collapsed MMP and rdCL, having normal morphology according to FSC and SSC (Fig. 3a). This means that more than 85% of cells were viable (Q2 + Q4), with a parallel decrease in early live-apoptotic cells (Q3) and dead cells (Q1) (Fig. 3a, b). The collapse of MMP with CCCP was either high, as it occurred at 10 μM, or low as with 5 μM, and yet in both situations the viability rose to about 90% (Fig. 3b). Therefore, MMP was needless for survival under hypoxia.

To complement this manual flow cytometry quadrant analysis using bivariate plots, we conducted an unsupervised classification through the FlowSOM algorithm [29], and we visualized the five dimensional phenotypic space (FSC, SSC, NAO, TMRE and PI) using tSNE plots [30]. This multidimensional analysis was in agreement with the manual analysis, and even detected higher values of live cells rescued from hypoxia-induced cell death by CCCP treatment over the 5–10 μM dose range. In addition, the study confirmed that rescued cells had either high MMP or low MMP, depending on the CCCP dose. At 5 μM CCCP dose, rescued cells showed the highest proportion in the high MMP group (matching Q2), while at 10 μM CCCP rescued cells were all in the low MMP group (matching Q4). In both cases rescued cells showed high NAO fluorescent signal (rdCL content). Early alive apoptotic cells matched the Q3 population in quadrant analysis, and were detected at 3 μM CCCP or below, almost disappearing at higher CCCP doses (Supp. Fig. S1).

The MM was reduced within the dose range, where CCCP rescued BC-K562 cells from hypoxia-induced damage (Fig. 3b). Conversely, CCCP reduced glucose uptake within the dose range where cell survival was increased (4-10 μM), even approaching the baseline values observed in normoxia (Fig. 3c).
3.4 ATO had a similar cytotoxic potency in normoxia and hypoxia

The dose–response curve of ATO and its EC50 under hypoxia was similar to the one obtained in normoxia (10.4 μM and 9.63 μM respectively) (Figs. 1b, 3g). However, we must consider that untreated cells had more than 50% death (red dots on the y-axis in Fig. 3g), meaning that at low doses ATO also had a rescuing effect (between 1 and 5 μM). Above EC50 cell death was increased in a dose-dependent manner (Fig. 3d, e, g). Contrasting CCCP results, there were no cells with collapse of MMP and rdCL (Q4) (Fig. 3e). Glucose uptake was reduced within the low dose range of ATO, although it attained the same uptake values of untreated cells at EC50 dose, well above the values observed in normoxia. (Fig. 3f).

3.5 Inhibition of mitophagy with Mdivi1 or VCR increased MM and rescued BC-K562 cells from hypoxia-induced death

Low performing mitochondria are pruned by fission and cannot undergo further fusion, but are rather transported within mitophagosomes along microtubules to the lysosomes (mitophagy). Therefore, we blocked mitophagy upstream with the fission inhibitor Mdivi1, and downstream with the microtubules transport inhibitor VCR, assuming that blocking mitophagy would be detrimental for BC-K562 cell survival under hypoxia. Contrary to our expectations, both drugs rescued BC-K562 cells from hypoxia-induced cell death. Viability was increased up to 80% under hypoxia with VCR concentrations between 1 and 6 μM (Fig. 4a, b). VCR increased MM as expected from blocking the elimination of mitochondria (Fig. 4b). In addition, cells maintained high MMP (Fig. 4c). Blocking mitophagy with Mdivi1 rescued cells from hypoxia-induced death, with viability figures close to 70% (Fig. 4d). This effect was only observed below 50 μM and higher doses of Mdivi1 were cytotoxic (Fig. 4e, f). However, at 50 μM Mdivi1 there was a very large increase in MM, something that also negates the classical view that mitochondria must be eliminated to improve tolerance to hypoxia (Fig. 4g).

3.6 VPA and DCA increased MM and rescued BC-K562 cells from hypoxia-induced death

VPA induces genes involved in mitochondrial biogenesis (PGC1α, TFAM) and we assumed VPA-treatment of BC-K562 cells would be deleterious to survival under hypoxia. However, VPA rescued cells from hypoxia-induced death, particularly within the dose range 1–2 mM, with figures even above 90% of viability (Fig. 5a, b). The cells showed high MMP and high MM (Fig. 5b). On the other hand, this increase in MM under hypoxia was paradoxically associated to a decrease in glucose uptake in VPA-treated cells compared to untreated cells. Even at high doses of VPA with a large increase in MM and decreased survival rates, glucose uptake remained low (Fig. 5c). This result suggested a connection between high MM and alternative metabolic sources to sustain glycolysis, such as biosynthetic metabolic pathways that intersect at the mitochondria.

DCA inhibits PDK1 and increases PDH activity. Therefore, it fosters entrance of pyruvate to the Krebs cycle and formation of acetyl-CoA and citrate. This effect is opposite to the action of HIF that increases expression of PDK1 and blocks PDH. By improving Krebs cycle and OXPHOS, DCA is considered a drug with an “anti-Warburg effect” that would be highly toxic under hypoxia.

However, BC-K562 were rescued from hypoxia-induced death by 1–3 mM DCA with viability above 80% (Fig. 5d, e, f). Unexpectedly, there was a dose-dependent increase in MM. Therefore, DCA had two HIF-opposing effects: increased MM instead of reduction by mitophagy, and increased PDH activity, yet these effects resulted in rescue of cells from hypoxia-induced death. Moreover, DCA decreased glucose uptake (Fig. 5g), again pointing to mitochondrial anabolic pathways supporting glycolysis in an unconventional form. The beneficial effect of increased MM by 3 mM DCA appeared to be independent of OXPHOS, since it was completely preserved when combined to CCCP over a range 1–10 μM (Supp. Fig. S2). Contrasting the effect of single drugs, the VPA + VCR combination was synergistic.
and highly cytotoxic for survival under hypoxia (Supp. Fig. S3a, b). We observed very large dose reduction indexes, especially for VCR. Therefore, blockage of mitophagy with increased mitochondrial biogenesis appeared to break the delicate balance between mitochondrial quality and quantity.
3.7 Genes differentially expressed in BC-K562 cells under hypoxia

All drugs tested had paradoxical responses under hypoxia, with most of them having patent rescuing effects from hypoxia-induced death instead of cytotoxicity. Since these drugs target mitochondria or MRCSR, paradoxical effects could be related to hypoxia-induced metabolic changes. In a recent study, Jain et al., exposed BC-K562 to 1% O₂ and 21% O₂ during 72 h, performed RNAseq on three samples under each condition, and deposited raw data in the curated GEO database (GSE144527) [32]. Since experimental conditions and time of exposure matched our own study with BC-K562 cells, we used this RNAseq dataset to explore MP changes induced by hypoxia from a gene expression perspective.

The analysis of DEG through the R package limma, showed significantly increased expression of genes involved in HIF stabilization (P4HA1, P4HB), mitophagy (BNIP3, BNIP3L, SQSTM1, ATG13, MAP1LC3B), increased glucose uptake (SLC2A3/GLUT3), regulation of glycolysis and pentose phosphate pathway (PKM1/2, LDHA, HK2), cholesterol synthesis (HMGT, HMGCS1, MVN), and fatty acid synthesis (FASN). The malic enzyme (ME), and the anti-apoptotic protein MCL1 were also upregulated, while many genes involved in anti-oxidant response were down regulated under hypoxia including CAT, PRDX1, PRDX2, PRDX4, and GSS. Also, MFN2 and OPA1, involved in mitochondrial fusion, were downregulated under hypoxia (Supp. Table S1). Analysis through Gene Ontology (GO) knowledge-base, conducted using the GOANA function of limma R package, showed GO terms with significantly DEG under hypoxia that were associated to mitochondrial cellular components, metabolism-related biological processes and functions (Supp. Table S2). The analysis using Kegg database through KEGGA function of limma R package, showed significantly DEG in pathways like glycolysis and gluconeogenesis, NADH regeneration, HIF-1 signalling, autophagy and steroid biosynthesis among others (Supp. Table S3).

Gene set enrichment analysis (GSEA) showed significantly enriched Hallmark gene sets under hypoxia that included “Hypoxia response” (p < 0.001; FDR < 0.001), “Glycolysis” (p < 0.001; FDR < 0.007), “Cholesterol Homeostasis” (p < 0.001; FDR < 0.001), “Apoptosis” (p < 0.001; FDR < 0.001), “MTORC1 signalling” (p < 0.001; FDR < 0.002), “KRAS signalling UP” (p < 0.008; FDR < 0.004), and “Epithelial mesenchymal transition” (p < 0.001; FDR < 0.001) (Supp. Table S4, Fig. 6a). Conversely, among significantly enriched Hallmark gene sets under normoxia we noticed “MYC targets” (p < 0.001; FDR < 0.001) and “Oxidative Phosphorylation” (p < 0.001; FDR < 0.001) (Supp. Table S5, Fig. 6a).

3.8 A metabolic plasticity signature derived from DEG and GSEA analysis

Considering leading edge genes from GSEA and significantly DEG (Fig. 6a, Supp. Tables S1-Table S3), we chose 99 genes as a potential signature that appeared relevant to MP under hypoxia, including genes involved in OXPHOS and ETC, Krebs cycle, cholesterol synthesis, fatty acid synthesis, mitochondrial fusion, mitophagy, glycolysis, pentose phosphate pathway, antioxidant response, anti-apoptosis and alternative sources of biosynthetic precursors (Supp. Table S6). This MP-signature clearly discriminated between BC-K562 cells under hypoxia and normoxia (Supp. Fig. S4a, Table S7).

3.9 Evaluation of the MP-signature in a transcriptomic dataset from CML patients

We further explored if this MP-signature could also discriminate between BC-CML patients and CP-CML patients, to ascertain if genes involved in MR along the disease progression, could match genes involved in MP (correspondence between the two metabolic dimensions). To this end, we explored the MP-signature among cells from CML patients samples in BC, AP, and CP, by using the GSE4170 microarray dataset. This dataset consisted of transcriptomes from a large CML patient’s cohort [33].

We conducted unsupervised classification of 113 patients based on their MP signature through hierarchical clustering, and we evaluated the first three most different groups. BC samples were classified mostly in one group, CP samples were classified in the two other groups, while AP samples were distributed among all three groups, but mainly with BC samples (Supp. Fig. S4b, c). Therefore, the MP-signature identified BC patients as a separated group, making evident a correspondence between MR and MP in the BC phenotype. The association between MP-signature category and clinical category was highly significant (p < 0.0001). When we next restricted the 99 gene set to a smaller signature of 25 genes (Supp. Table S8), 92% of BC cases corresponded to a single cluster (Cluster_2), and 95% CP cases corresponded to the other two clusters (Cluster_1 + 3). (Fig. 6b, c). About 46% of AP cases were in Cluster_2 together with BC cases, while the 53% remaining were in Cluster_1 + 3 with CP cases. Cluster_2 consisted of 77% BC cases and 16% AP cases (93% both), while Cluster_1 + 3 consisted of 84% CP cases (Fig. 6b, c). Noticeably, OXPHOS-related genes that were down-regulated
under hypoxia in BC-K562 cells, appeared characteristically up-regulated in BC patients compared to CP patients. By contrast, genes of cholesterol metabolism were up-regulated under hypoxia in BC-K562 cells, but were down-regulated when comparing BC samples to CP samples (Fig. 6c). This MP-signature made of 25 genes also distinguished BC-K562 cells under normoxia and hypoxia considering the 6 samples from GSE144527 (Fig. 6d). It is noticeably that most of these genes were unrelated to HIF-targets. The differential expression of many genes was dissimilar in the comparison between CP and BC (metabolic reprogramming), and the comparison between normoxia and hypoxia (metabolic plasticity). For example MFN2 was down-regulated in BC, and OPA1 was up-regulated in BC compared to CP. However, both MFN2 and OPA1 were down-regulated in BC-K562 cells under hypoxia compared to the same cells under normoxia. Therefore, the gene set that distinguished BC from CP was the same as the gene set that distinguished hypoxia from normoxia, but the sense and magnitude of differential gene expression in both situations was different. (Fig. 6c, d and Supp. Fig. S5). In fact there was an inverse correlation between differential expressions BC vs CP compared to hypoxia vs normoxia (Spearman rank correlation coefficient − 0.41; 95% CI [− 0.70, − 0.01], p < 0.05), (Fig. 6e). Therefore, the expression profile of this 25-gene set may reflect changes due to both MR and MP of blast crisis cells.

4 Discussion

In this work we explored the cytotoxic potency of drugs that target mitochondria and MRCSR under < 1% hypoxia during 72 h in BC-K562 cells. Less than 50% of untreated cells survive under these conditions. These figures are in complete agreement with a previous study that quantified dead and apoptotic cells at 72 h and 1% O2 [37]. In addition, further studies showed that surviving untreated-BC-K562 cells acquire traits of leukaemia stem cells at day seven, and even decrease Bcr-Abl expression at the protein level [38]. Paradoxically, many of the mitochondria-targeted drugs that we tested, rescued cells from hypoxia-induced death at 72 h instead of killing them. Since hypoxia represented a sudden environmental change driving rapid metabolic changes, we inferred that the cause of the profoundly altered cytotoxicity was linked to these changes, particularly those linked to mitochondrial functions, since mitochondria initiates apoptosis through MOMP. Mitochondria are not a significant source of ATP under hypoxia, and HIF-mediated response to hypoxia involves its elimination through mitophagy [10]. However, mitochondria may not be superfluous under hypoxia in BC cells, considering that abnormal anabolic pathways may interweave at the mitochondria [39]. The gene expression analysis of BC-K562 supports this idea, denoting high enrichment of gene sets associated to fatty acid synthesis and cholesterol homeostasis. These anabolic routes require a strong support of pathways operating at the mitochondria, such as citrate efflux and truncated forms of the Krebs cycle. How these anabolic rather than bioenergetic functions of mitochondria impact on apoptosis threshold is only partially known, and several hypothesis have been raised. Among these evidences are increased anti-apoptotic proteins such as MCL1, low performing ETC that is less prone to MOMP, and a stronger antioxidant capacity provided by large amounts of NADPH.

Cancer cells have several changes in metabolic pathways that increase along disease progression. These MR changes are considered to progress as waves of gene expression changes, occurring along extended periods of time [40]. However, cell metabolism is dependent on the microenvironment, and environmental conditions of cancer are not stable but rather often subject to sudden changes. Therefore, cancer cells may not only have an abnormal metabolism, but have also the capability of transiently changing the metabolic program [5]. This second dimension of transient metabolic changes are known as MP [41]. We inferred that MP would be greater in advanced disease, providing BC cells a faster metabolic adaptation to transient environmental changes such as hypoxia and lack of nutrients [42].

Our hypothesis was that the cause of paradoxical response to drugs under hypoxia could be related to metabolic changes of cells that could impact on drug targets. For example, ETC becoming disposable for ATP synthesis under hypoxia, but Krebs cycle still operating to support anabolic pathways, could transform a drug vulnerability under normoxia to a drug-induced growth advantage under hypoxia. To explore if such changes effectively occur, we compared
the transcriptomes of BC-K562 cells under normoxia and hypoxia, using raw data from an RNAsseq dataset that became available recently in GEO database (GSE144527) [32].

Our analysis of GSE144527 allowed us to define a MP-signature of 25 genes related to mitochondrial function, mitophagy, mitochondrial fusion, Krebs cycle enzymes, glycolysis, fatty acid synthesis, and cholesterol metabolisms that were differentially expressed in BC-K562 after 72 h under 1% hypoxia. This MP-signature was sufficient to completely discriminate cells under normoxia and hypoxia through unsupervised classification, confirming that the occurrence of these metabolic changes was a distinctive feature of hypoxia-adaptation of cells, and could potentially impact on drug targets. Noticeably, this MP-signature had only a minor representation of classical HIF target genes. This shows how critical and archetypal is this expression profile in hypoxia adaptation of BC-K562 cells.

In agreement with a prominent role of mitochondrial genes under hypoxia, we found that increased MM was associated with rescue from hypoxia-induced death. This increase resulted from very dissimilar grounds such as blocking mitophagy (VCR and Mdivi1), increased mitochondrial biogenesis (VPA), and even increased MM by DCA probably owing...
Fig. 6 A 25-gene metabolic plasticity signature derived from BC-K562 cells under hypoxia. a Gene set enrichment analysis of BC-K562 cells exposed to 1% hypoxia for 72 h. The RNA expression values of 4362 genes of three replicates under hypoxia and three replicates under normoxia were compared over 50 functional Hallmark gene lists (MSigDB). Enrichment, indicating RNA over or under-expression, is represented by profile line graphs, where the black bar represents a single gene from the list of 4362 genes, which matched a gene of the Hallmark list (hit), and the line represents over-expression (left peak) or under-expression (right peak) in a comparison hypoxia vs normoxia, or normoxia vs hypoxia (hypoxia-enriched, and normoxia-enriched respectively). Clustered black bars on the left indicate over-expression, and clustered black bars on the right indicate under-expression. The enrichment score (ES) provides a magnitude of over-expression (positive value) or under-expression (negative value) of genes of the Hallmark functions (see Materials and Methods section). The detailed genes (hits) found in each Hallmark gene list are shown as heatmaps on the right (they correspond to the black bars in the enrichment profile graphs shown on the left). Red squares are genes over-expressed and blue squares are gene under-expressed in hypoxia. Each column of the heatmaps represents an RNAseq sample. Since the middle part shows genes expressed similarly in hypoxia and normoxia we omitted the middle part (indicated by three large dots). b Contingency table exploring association between clinical categories CP, AP and BC and unsupervised classification of 113 CML patients according to their RNA expression profile. The patients were classified into Cluster_1, Cluster_2 and Cluster_3, considering only RNA expression values of a 25-gene metabolic plasticity signature, selected among those genes that we found differentially expressed in BC-K562 cells under hypoxia. Pie charts illustrates how the three clinical categories fit in two main “metabolic” clusters (Cluster_1 + 3 and Cluster_2) with CP and BC cases clearly separated, and AP cases fitting half in each cluster. c Heatmap showing expression profile with unsupervised classification of 113 CML patients using the 25-gene metabolic plasticity signature obtained from BC-K562 cells under hypoxia. The 25 genes are listed on the left of the heatmap. BC patients are clearly associated to the profile of cluster_2 while CP patients are associated to profiles of Cluster_1 + 3. d Heatmap presenting unsupervised classification of six BC-K562 transcriptomes and showing that the 25-gene MP-signature also discriminates between BC-K562 cells under hypoxia and normoxia. e Correlation plot showing that the gene expression profile of these 25 genes is different and inversely correlated when comparing BC patients to CP patients (metabolic reprogramming) than when comparing BC-K562 cells under hypoxia vs normoxia (metabolic plasticity).
to targeting PGC1α as off-target, apart from its main action increasing PDH activity (we provide the analysis of a microarray dataset of BC-K562 cells treated with 1 mM VPA that was available from GEO database in Supp. Fig. S6a).

The exception was CCCP that decreased MM while rescuing cells from hypoxia-induced cell death. This may be due to its massive induction of mitophagy. In fact, CCCP is used as a positive control of mitophagy in experimental settings [43]. The beneficial effect of increased MM was due to augmented ETC and OXPHOS since this is hampered by low O2. In fact, CCCP caused rescue from hypoxia-induced death while completely preventing the function of ETC.

Moreover, CCCP + DCA rescued cells with complete collapse of MMP and increased MM (particularly at the highest dose of 10 μM CCCP), enforcing the idea that the beneficial effect of increased MM was other than ETC and OXPHOS (Supp. Fig. S2). It is important to emphasize that collapse of MMP with rdCL content (high NAO fluorescence signal) does not indicate mitochondrial damage, but just cells where there is no possibility of OXPHOS, because without MMP the ETC becomes non-functional. On the other side, CCCP does not compromise the Krebs cycle and allows other mitochondrial anabolic pathways to continue fulfilling their functions. The anabolic functions of mitochondria involve an active Krebs cycle with cataplerotic pathways such as citrate or malate efflux, inter-conversion with amino acids such as aspartic acid (oxaloacetate) and alanine (pyruvate), and continuous replenishment through anaplerotic pathways such as glutaminolysis (Fig. 7) [3]. The increase in PDH activity as caused by DCA fosters citrate formation from acetylCoA and oxalacetate. Therefore, the outstanding pro-survival effect of DCA can be linked to citrate efflux and support of biosynthetic pathways in the cytoplasm, such as cholesterol homeostasis and fatty acid synthesis, with high anti-oxidant capacity provided by increased NADPH (Fig. 7). In contrast to DCA, ATO inhibits PDH and 2OGDH [17]. These two enzymes are critical for Krebs cycle, even when it operates only in support of anabolic pathways. Therefore, ATO targets the core of both bioenergetic and anabolic functions of mitochondria (we provide the analysis of a microarray dataset of BC-K562 cells treated with 1.5 μM ATO that was available from GEO database in Supp. Fig. S6b). Krebs cycle is required for anoxic glutaminolysis, where citrate is formed by reductive carboxylation (reverse and truncated Krebs cycle), and is also required in forward-glutaminolysis where a truncated Krebs cycle is coupled to ETC complex III through succinate, and allows citrate efflux by combination of oxalacetate and acetyl CoA (Fig. 7) [3, 44, 45]. Therefore, it is not surprising that ATO was the only toxic drug in hypoxia for BC-K562 cells, hampering all forms of mitochondrial functions. In addition, increased MM with lack of mitochondrial quality control provided by blockage of mitophagy, was highly toxic showing exceedingly high synergism and dose reduction indexes (Supp. Fig. S3). This result implies that the benefit provided by increased MM is abruptly offset beyond a certain level of accumulated mitochondrial defects.

Noticeably, survival from hypoxia-induced death was associated not only with increased MM but with decreased glucose uptake compared to untreated cells, and even approaching values comparable to normoxia. This is a remarkable result, since ATP must be provided by increased glycolysis under hypoxia. The DEG and GSEA analysis confirmed increased glycolysis in untreated cells. However, we observed a consistent decrease of glucose uptake in drug-rescued surviving cells. This suggests that glycolysis may be somehow aided by anabolic pathways fostered by mitochondria through processes such as citrate efflux (Fig. 7) [3]. After GU and conversion of glucose into glucose-6-phosphate (G6P), glucose carbons may take a number of alternative downstream routes, including glycolysis and ATP production, and diversion into the PPP pathway for the supply of biosynthetic precursors and NADPH [46]. It is possible that increased availability of mitochondria-derived precursor metabolites for biosynthesis processes could decrease the need for routing of G6P to PPP, increasing the availability for the glycolysis pathway. In this way, the demand for GU and conversion to G6P could be decreased to some extent. The regulation of these metabolic pathways appears highly dynamic and complex, and it is far from being understood in cancer cells, although some studies propose a role for isoforms of PKM in controlling the balance between G6P routed to PPP and to glycolysis according to the bioenergetic and biosynthetic demands [47, 48]. In this sense, we could speculate that cytoplasmic and mitochondrial metabolic pathways would be coordinated in such a way that, the more mitochondria supporting anabolic roles, the less the demand for GU given a certain level of demand for bioenergetic and biosynthesis processes.

The paradoxical effects of mitochondria-targeted drugs with increased MM and decreased glucose uptake occurred in cells that without treatment showed changes in the 25-gene MP signature. Since MP represents a transient adaptation driven by the environment, we asked if these same gene set could work as a MR-signature in samples from CML patients. Blast crisis cells showed a different expression profile in this 25-gene set, and this MR-signature discriminated between samples from BC patients and CP patients. Therefore, the expression profile of the same gene set could capture changes between hypoxia and normoxia, and changes between CP and BC patients. However, this profile in BC cells was different in both comparisons. In fact, the expressions of these genes were inversely correlated. Genes that were downregulated in response to hypoxic environment, were often those upregulated in BC patients compared to cells in CP patients. This again underscores that MR is an average profile correlated to progression, and that expression changes may occur in a
Fig. 7 A simplified diagram representing some relevant anabolic and catabolic pathways intersecting at mitochondria that may be relevant in metabolic plasticity of BC-K562 cells and the paradoxical effects of drugs under hypoxia. The Krebs cycle can operate as a source of biosynthetic precursors through cataplerotic pathways such as citrate efflux, but must be compensated by anaplerotic pathways such as glutaminolysis as a source of 2-oxoglutarate (2-OG). A truncated Krebs cycle may operate in reverse sense even under complete absence of oxygen (anoxic glutaminolysis), or may operate in forward sense, generating FADH from succinate at ETC complex II, even with very low oxygen (forward Krebs glutaminolysis). However, forward Krebs glutaminolysis coupled to citrate efflux requires continuous supply of acetyl-CoA. This is inhibited by ATO and improved by DCA. In addition, ATO partially inhibits 2-oxoglutarate-dehydrogenase (2-OGDH). ROS-mediated damage to the ETC by ATO and CCCP may be reduced under lack of oxygen. The graph shows in light pink some genes significantly upregulated under hypoxia compared to normoxia (p-value < 0.05) in untreated BC-K562 cells (GSE144527), and how citrate efflux appears coupled to cholesterol and fatty acid synthesis. PPP pentose phosphate pathway, MCT1 monocarboxylate transporter1 /SLC16A1, HMG-CoA hydroxy-methyl-glutaryl-CoA, and the remaining are conventional gene symbols (https://www.genecards.org).
second metabolic dimension driven by environment (plasticity) that may involve not only O₂ but availability of nutrients such as glucose and amino acids [37]. Therefore, methods comparing two environments can unveil this second metabolic dimension and provide a better characterization of altered metabolism in advanced disease.

In a previous work, we found no correlation to disease progression of drug potencies in a single environment (normoxia or hypoxia), but a strong correlation to differential response of drugs targeting MRCSR between normoxia and hypoxia in CLL patients [25]. Here we substantiate those findings through gene expression profile, showing that an environment-driven differential signature was also highly correlated to progression in CML. We conclude that both MP and MR may impact drug potency. Understanding the interconnection of MP and MR with drug potency can give hints to overcome resistance. Recent studies have shown several successful examples including Venetoclax and ATO, TKIs and ATO, Venetoclax and CB839, and inhibitors of fatty acid oxidation such as Etomoxir, highlighting that the Krebs cycle appears as a relevant target to suppress anabolic functions of mitochondria [49–52].

Acknowledgements This work was supported by grant Universidad de Buenos Aires, Argentina (20720170100021BA), and grants CONICET PIP 11220170100146CO001 and in part by grant CONICET PUE 22920160100017CO

Author contributions LS: Investigation, data curation, visualization, methodology, formal analysis. MCL: conceptualization, writing—review and editing. MGF: investigation, data curation, visualization. ERR: writing—review and editing, funding acquisition, investigation. GAB: conceptualization, funding acquisition, writing—review and editing, data curation, investigation. All the authors read and approved the final manuscript.

Data availability The datasets generated during and/or analysed during the current study are available in the Gene Expression Omnibus (GEO) repository, [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE144527; https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE4170], are included in supplementary information files, and are available from the corresponding author on reasonable request.

Declarations

Competing interest The authors declare no conflicts of interest.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. Jabbour E, Kantarjian H. Chronic myeloid leukemia: 2020 update on diagnosis, therapy and monitoring. Am J Hematol. 2020;95(6):691–709. https://doi.org/10.1002/ajh.25792.
2. Radich JP. The biology of CML blast crisis. Hematology. 2007. https://doi.org/10.1182/asheducation-2007.1.384.
3. DeBerardinis RJ, Chandel NS. Fundamentals of cancer metabolism. Sci Adv. 2016;2(5): e1600200. https://doi.org/10.1126/sciadv.1600200.
4. Noel BM, Ouellette SB, Marholz L, Dickey D, Navis C, Yang TY, et al. Multiomic profiling of tyrosine kinase inhibitor-resistant K562 cells suggests metabolic reprogramming to promote cell survival. J Proteome Res. 2019;18(4):1842–56. https://doi.org/10.1021/acs.jproteome.9b00028.
5. Koczula KM, Ludwig C, Hayden R, Cronin L, Pratt G, Parry H, et al. Metabolic plasticity in CLL: adaptation to the hypoxic niche. Leukemia. 2016;30(1):65–73. https://doi.org/10.1038/leu.2015.187.
6. Porporato PE, Filigheddu N, Pedro JMB, Kroemer G, Galluzzi L. Mitochondrial metabolism and cancer. Cell Res. 2018;28(3):265–80. https://doi.org/10.1038/cr.2017.155.
7. Bock FJ, Tait SWG. Mitochondria as multifaceted regulators of cell death. Nat Rev Mol Cell Biol. 2020;21(2):85–100. https://doi.org/10.1038/s41580-019-0173-8.
8. Sharma A, Boise LH, Shanmugam M. Cancer metabolism and the evasion of apoptotic cell death. Cancers (Basel). 2019. https://doi.org/10.3390/cancers11081144.
9. Vidal RS, Quarti J, Rodrigues MF, Rumjanek FD, Rumjanek VM. Metabolic reprogramming during multidrug resistance in leukemias. Front Oncol. 2018;8:90. https://doi.org/10.3389/fonc.2018.00090.
10. Semenza GL. Regulation of oxygen homeostasis by hypoxia-inducible factor 1. Physiology (Bethesda). 2009;24:97–106. https://doi.org/10.1152/physiol.00045.2008.
11. Zhang H, Bosch-Marce M, Shimoda LA, Tan YS, Baek JH, Wesley J, et al. Mitochondrial autophagy is an HIF-1-dependent adaptive metabolic response to hypoxia. J Biol Chem. 2008;283(16):10892–903. https://doi.org/10.1074/jbc.M800102200.

12. Cavaliere V, Lombardo T, Costantino SN, Kornblihtt L, Alvarez EM, Blanco GA. Synergism of arsenic trioxide and MG132 in Raji cells attained by targeting BNIP3, autophagy, and mitochondria with low doses of valproic acid and vincristine. Eur J Cancer. 2014;50(18):3243–61. https://doi.org/10.1016/j.ejca.2014.09.012.

13. Kornblihtt L, Carreras M, Blanco G. Targeting mitophagy in combined therapies of haematological malignancies. In: Gorbunov N, editor. Discover Oncology (2022) 13:60 | https://doi.org/10.1007/s12672-022-00524-y

14. de Graaf AO, van den Heuvel LP, Dijkman HB, de Abreu RA, Birkenkamp KU, de Witte T, et al. Bcl-2 prevents loss of mitochondria in CCCP-induced apoptosis. Exp Cell Res. 2004;299(2):533–40. https://doi.org/10.1016/j.yexcr.2004.06.024.

15. Lombardo T, Folgar MG, Salaverry L, Rey-Roldan E, Alvarez EM, Carreras MC, et al. Regulated cell death of lymphoma cells after graded mitochondrial damage is differentially affected by drugs targeting cell stress responses. Basic Clin Pharmacol Toxicol. 2018;122(5):489–500. https://doi.org/10.1111/bcpt.12945.

16. Miller WH Jr, Schipper HM, Lee JS, Singer J, Waxman S. Mechanisms of action of arsenic trioxide. Cancer Res. 2002;62(14):3893–903.

17. Bergquist ER, Fischer RJ, Martin BD. Inhibition by methylated organo-arsenicals of the respiratory 2-oxo-acid dehydrogenase complex. J Organomet Chem. 2009;694(6):973–80. https://doi.org/10.1016/j.jorganchem.2008.12.028.

18. Luo X, Liu R, Zhang Z, Chen Z, He J, Liu Y. Mitochondrial division inhibitor 1 attenuates mitophagy in a rat model of acute lung injury. Biomed Res Int. 2019;2019:2193706. https://doi.org/10.1155/2019/2193706.

19. Twig G, Erorza A, Molina AJ, Mohamed H, Wikström JD, Walzer G, et al. Fission and selective fusion govern mitochondrial segregation and elimination by autophagy. EMBO J. 2008;27(2):433–46. https://doi.org/10.1038/sj.emboj.6200763.

20. Mackeh R, Perdiz D, Sorin L, Codogno P, Pous C. Autophagy and microtubules—new story, old players. J Cell Sci. 2013;126(5):1071–80. https://doi.org/10.1242/jcs.115626.

21. Sitarz KS, Elliott HR, Karaman BS, Reton C, Chinnery PF, Horvath R. Valproic acid triggers increased mitochondrial biogenesis in POG-deficient fibroblasts. Mol Gen Metab. 2014;112(1):57–63. https://doi.org/10.1016/j.mgen.2014.03.006.

22. Cook KM, Shen H, McKelvey KJ, Gee HE, Hau E. Targeting glucose metabolism of cancer cells with dichloroacetate to radiosensitize high-grade gliomas. Int J Mol Sci. 2021. https://doi.org/10.3390/ijms22147265.

23. Lozio CB, Lazzio BB. Human chronic myelogenous leukemia cell-line with positive Philadelphia chromosome. Blood. 1975;43(3):321–34.

24. Weinstein JN, Pommier Y. Transcriptomic analysis of the NCI-60 cancer cell lines. C R Biol. 2003;326(10–11):909–20. https://doi.org/10.1016/j.crvi.2003.08.005.

25. Kornblihtt LI, Cabral Lorenzo MC, Gil Folgar ML, Minissale CJ, Malusardi C, Rojas F, et al. A drug potency signature links progression of chronic lymphocytic leukemia to mitochondria-related stress responses and metabolic reprogramming under hypoxia. Toxicol Appl Pharmacol. 2020;398:115016. https://doi.org/10.1016/j.taap.2020.115016.

26. Wu D, Yotnda P. Induction and testing of hypoxia in cell culture. J Vis Exp. 2011. https://doi.org/10.3791/2899.

27. Mathupala SP, Kioussis S, Szerlip NJ. A lab assembled microcontroller-based sensor module for continuous oxygen measurement in portable hypoxia chambers. PLoS ONE. 2016;11(2):e0148923. https://doi.org/10.1371/journal.pone.0148923.

28. Li XX, Tsoi B, Li YF, Kurihara H, He RR. Cardiolipin and its different properties in mitophagy and apoptosis. J Histochem Cytochem. 2015;63(5):301–11. https://doi.org/10.1369/jjhc.2015.541577.

29. Van Gassen S, Cellebaut B, Van Helden MJ, Lambrecht BN, Demeester P, Dhaene T, et al. FlowSOM: Using self-organizing maps for visualization and interpretation of cytometry data. Cytometry A. 2018;87(7):636–45. https://doi.org/10.1002/cyto.a.22625.

30. Belkina AC, Ciccolella CO, Anno R, Halpert R, Spidlen J, Snyder-Cappione JE. Automated optimized parameters for T-distributed stochastic neighbor embedding improve visualization and analysis of large datasets. Nat Commun. 2019;10(1):5415. https://doi.org/10.1038/s41467-019-13055-y.

31. Chou TC. Drug combination studies and their synergy quantification using the Chou-Talalay method. Cancer Res. 2010;70(2):440–6. https://doi.org/10.1158/0008-5472.CAN-09-1947.

32. Jain IH, Calvo SE, Markhard AL, Skinner OS, To TL, Ast T, et al. Genetic screen for cell fitness in high or low oxygen highlights mitochondrial and lipid metabolism. Cell. 2020;181(3):716-727 e11. https://doi.org/10.1016/j.cell.2020.03.029.

33. Radich JP, Dai H, Mao M, Oehler V, Schelter J, Druker B, et al. Gene expression changes associated with progression and response in chronic myeloid leukemia. Proc Natl Acad Sci USA. 2006;103(8):2794–9. https://doi.org/10.1073/pnas.0510423103.

34. Ritchie ME, Phipson B, Wu D, Hu Y, Law CW, Shi W, et al. limma powers differential expression analyses for RNA-sequencing and microarray studies. Nucleic Acids Res. 2015;43(7):e47. https://doi.org/10.1093/nar/gkv007.

35. Subramanian A, Tamayo P, Mootha VK, Mukherjee S, Ebert BL, Gillette MA, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci USA. 2005;102(43):15545–50. https://doi.org/10.1073/pnas.0506580102.

36. Kassambara AU. Practical Guide to Cluster Analysis in R: Unsupervised Machine Learning. STHDA: New York; 2017.

37. Giuntoli S, Tanturi M, Di Gesualdo F, Barbetti V, Rovida E, Dello Sbarba P. Glucose availability in hypoxia regulates the selection of chronic myeloid leukemia progenitor subsets with different resistance to imatinib-mesylate. Haematologica. 2011;96(2):204–12. https://doi.org/10.3324/haematol.2010.029082.

38. Tanturi M, Giuntoli S, Barbetti V, Rovida E, Dello Sbarba P. Hypoxia selects bortezomib-resistant stem cells of chronic myeloid leukemia. PLoS ONE. 2011;6(2):e17008. https://doi.org/10.1371/journal.pone.0017008.

39. Oliveira GL, Coelho AR, Marques R, Oliveira PJ. Cancer cell metabolism: rewiring the mitochondrial hub. Biochim Biophys Acta Mol Basis Dis. 2021;201672(6):166016. https://doi.org/10.1016/j.bbadis.2020.166016.

40. Smolkova K, Plecita-Hlavata L, Bellance N, Benard G, Rossignol R, Jezeck P. Waves of gene regulation suppress and then restore oxidative phosphorylation in cancer cells. Int J Biochem Cell Biol. 2011;43(7):950–68. https://doi.org/10.1016/j.biocel.2010.05.003.

41. Shuvakov O, Daks A, Fedorova O, Petukhov A, Barlev N. Linking metabolic reprogramming, plasticity and tumor progression. Cancers (Basel). 2021. https://doi.org/10.3390/cancers13040762.
42. McGuirk S, Audet-Delage Y, St-Pierre J. Metabolic fitness and plasticity in cancer progression. Trends Cancer. 2020;6(1):49–61. https://doi.org/10.1016/j.trecan.2019.11.009.

43. Narendra D, Tanaka A, Suen DF, Youle RJ. Parkin is recruited selectively to impaired mitochondria and promotes their autophagy. J Cell Biol. 2008;183(5):795–803. https://doi.org/10.1083/jcb.200809125.

44. Le A, Lane AN, Hamaker M, Bose S, Gouw A, Barbi J, et al. Glucose-independent glutamine metabolism via TCA cycling for proliferation and survival in B cells. Cell Metab. 2012;15(1):110–21. https://doi.org/10.1016/j.cmet.2011.12.009.

45. Smolikova K, Jezeck P. The role of mitochondrial NADPH-dependent isocitrate dehydrogenase in cancer cells. Int J Cell Biol. 2012;2012:273947. https://doi.org/10.1155/2012/273947.

46. Shulman RG, Rothman DL. The glycogen shunt maintains glycolytic homeostasis and the warburg effect in cancer. Trends Cancer. 2017;3(11):761–7. https://doi.org/10.1016/j.trecan.2017.09.007.

47. Yang GJ, Wu J, Leung CH, Ma DL, Chen J. A review on the emerging roles of pyruvate kinase M2 in anti-leukemia therapy. Int J Biol Macromol. 2021;193(Pt B):1499–506. https://doi.org/10.1016/j.ijbiomac.2021.10.213.

48. Zhang Z, Deng X, Liu Y, Liu Y, Sun L, Chen F. PKM2, function and expression and regulation. Cell Biosci. 2019;9:52. https://doi.org/10.1186/s13578-019-0317-8.

49. Cho H, Jang JE, Eom JJ, Jeung HK, Chung H, Kim JS, et al. Arsenic trioxide synergistically promotes the antileukaemic activity of venetoclax by downregulating Mcl-1 in acute myeloid leukaemia cells. Exp Hematol Oncol. 2021;10(1):28. https://doi.org/10.1186/s40164-021-00221-6.

50. Estan MC, Calvino E, Calvo S, Guillen-Guio B, Boyano-Adanez Mdel C, de Blas E, et al. Apoptotic efficacy of etomoxir in human acute myeloid leukemia cells. Cooperation with arsenic trioxide and glycolytic inhibitors, and regulation by oxidative stress and protein kinase activities. PLoS ONE. 2014;9(12):e115250. https://doi.org/10.1371/journal.pone.0115250.

51. Jacque N, Ronchetti AM, Larrue C, Meunier C, Birsen R, Willems L, et al. Targeting glutaminolysis has antileukemic activity in acute myeloid leukemia and synergizes with BCL-2 inhibition. Blood. 2015;126(11):1346–56. https://doi.org/10.1182/blood-2015-01-621870.

52. Zhu HH, Qian JJ, Sun WJ, You LS, Wang QQ, Naranmandura H, et al. Venetoclax and arsenic showed synergistic anti-leukemia activity in vitro and in vivo for acute myeloid leukemia with the NPM1 mutation. Am J Hematol. 2020;95(3):ES5–7. https://doi.org/10.1002/ajh.25719.

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.