Quantifying small molecule phenotypic effects using mitochondrial morpho-functional fingerprinting and machine learning

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In primary fibroblasts from Leigh Syndrome (LS) patients, isolated mitochondrial complex I deficiency is associated with increased reactive oxygen species levels and mitochondrial morpho-functional changes. Empirical evidence suggests these aberrations constitute linked therapeutic targets for small chemical molecules. However, the latter generally induce multiple subtle effects, meaning that in vitro potency analysis or single-parameter high-throughput cell screening are of limited use to identify these molecules. We combine automated image quantification and artificial intelligence to discriminate between primary fibroblasts of a healthy individual and a LS patient based upon their mitochondrial morpho-functional phenotype. We then evaluate the effects of newly developed Trolox variants in LS patient cells. This revealed that Trolox ornithylamide hydrochloride best counterbalanced mitochondrial morpho-functional aberrations, effectively scavenged ROS and increased the maximal activity of mitochondrial complexes I, IV and citrate synthase. Our results suggest that Trolox-derived antioxidants are promising candidates in therapy development for human mitochondrial disorders.

Virtually every cell contains mitochondria, which are double membrane organelles that play multiple roles in cell metabolism, signal transduction and ATP generation. Mitochondrial function is sustained by the action of the oxidative phosphorylation (OXPHOS) system, consisting of four electron transport chain (ETC) complexes (CI-CIV) and the ATP-generating F1F0-ATPase (CV). ETC and CV action is linked via the highly negative potential (ΔΨ) across the mitochondrial inner membrane (MIM) through chemiosmotic coupling. Malfunction of the OXPHOS system is observed in many human diseases including metabolic syndromes such as Leigh Syndrome (LS)1–3. At the cellular level mitochondrial dysfunction and alterations in mitochondrial structure and ΔΨ are often paralleled by increased reactive oxygen species (ROS) levels4–9. This suggests that these aberrations might constitute linked therapeutic targets. In fibroblasts from LS patients with isolated CI deficiency (OMIM 253010), the α-tocopherol derivative Trolox (6-hydroxy-2, 5, 7, 8-tetramethylchroman-2-carboxylic acid) normalized increased levels of CM-H2DCF (5-(and-6)-chloromethyl-2′,7′-dichlorodihydro-fluorescein) oxidizing ROS10. Trolox treatment also increased the amount/activity of the CI holo complex and improved Ca2+ stimulated mitochondrial ATP production in LS cells10,11. In fibroblasts from a healthy individual (CT) Trolox stimulated mitochondrial length and degree of branching12. Also studies in other models of mitochondrial disease highlighted beneficial effects of (mitochondria-targeted) antioxidants13–16. Although the above suggests that elevated ROS levels might play a pathophysiological role in mitochondrial dysfunction, antioxidant treatment of “real” mitochondrial disease patients appeared only (transiently) effective in a very limited number of cases17–19. These negative results raised doubts about the effectiveness of antioxidant treatment in ROS-related human pathologies20–24.
The discrepancy between results obtained in model systems and mitochondrial disease patients is striking and might be explained by the dual role of ROS as signaling and damaging molecules. In addition, antioxidants can also display pro-oxidant properties depending on their concentration, physicochemical properties and reaction environment. Importantly, therapeutic small molecules (including bioactive antioxidants) generally target multiple effectors thereby exerting subtle pleiotropic effects (both on- and off-target) at the cellular level. This means that simply studying the in vitro potency and/or performing single-parameter high-throughput image-based cell screening is of limited use to identify potential drug-like molecules or understand their mode-of-action. Therefore the rate-of-success during lead selection and optimization in drug discovery benefits from multi-parameter phenotypic profiling.

Here we present an integrated experimental and computational strategy that is broadly applicable for small molecule profiling and based on quantifying their phenotypical effects. This approach consists of: (i) automated calculation of phenotypic “fingerprints” reflecting mitochondrial morphology and ΔΨ (“morpho-function”) at the level of individual mitochondria in single living cells, (ii) using these morpho-functional fingerprints for supervised machine learning classification of CT and LS patient cells, (iii) automated evaluation of morpho-functional effects induced by small molecule oxidants in LS patient cells. In a proof-of-principle study, the above strategy was applied to evaluate the phenotypic effects of four newly discovered Trolox variants in LS patient cells. This highlighted Trolox ornithylamide hydrochloride (KH003) as a therapeutically promising Trolox derivative, as confirmed in vitro in cellulo analysis. KH003 displayed cellular ROS scavenging, stimulated CI, CIV and citrate synthase activity. This suggests that Trolox-derivatives are promising candidates in therapy development for human mitochondrial disorders.

**Results**

Mitochondrial morpho-functional fingerprinting of human skin fibroblasts. An automated approach was developed for phenotypic analysis of mitochondrial morphology and ΔΨ (“morpho-functional fingerprinting”) in primary human skin fibroblasts (Fig. 1). Cells were stained with TMRM (tetramethyl rhodamine methyl ester), a mitochondria-specific cation that accumulates in the mitochondrial matrix in a ΔΨ-dependent manner, and visualized using epifluorescence microscopy (Fig. 1A; yellow boxes). Various quantitative parameters (Supplementary Table S1) describing mitochondrial morphology and TMRM intensity (“morpho-functional descriptors”) were extracted from the microscopy images by applying an automated image processing and analysis algorithm (Fig. 1A; blue boxes). This strategy was extensively validated previously in primary human skin fibroblasts (Supplementary Information). For every image, 31 descriptors were calculated for each mitochondrial object. Relative to typical control cells (CTS120), descriptor values in LS patient cells (P5175) were unaffected (8 descriptors), significantly increased (2 descriptors) or significantly decreased (21 descriptors; Supplementary Table S1).

Using mitochondrial morpho-functional fingerprints for supervised machine learning and cell classification. The dataset of 31 descriptors was used for training a machine learning algorithm to discriminate between CTS120 and P5175 cells. In total, 711 images were analyzed for the CTS120 cell line (containing 187465 mitochondrial objects) and 567 images were analyzed for the P5175 cell line (112615 objects). By calculating the median value of the 31 descriptors per image, a morpho-functional “fingerprint” of the P5175 cell line was obtained (Supplementary Table S1). Using the median value instead of the average value prevents that outlier values bias the analysis. Next, we used the multivariate fingerprints to determine whether various supervised machine learning algorithms were able to reliably discriminate between CTS120 and P5175 cells (Fig. 1A; green boxes). For this purpose, the data set for the CTS120 and P5175 cells was equally divided into a “training set” and a “test set”. The training set was used to “teach” the multivariate machine learning model, after which the test set was used to assess its classification (predictive) performance. Crucially, the test images were not used to train the model.
Figure 2 | Machine learning classification of control and patient cells. (A) Visualization of the test samples according to the logistic regression scores vs. the main source of variance in the data (Principal component 1: PC1). Data from individual microscopy images of control cells (CT5120; green dots; n = 356 images; N = 13 days) and patient cells (P5175; red dots; n = 235 N = 13) separates along the horizontal axis. A cell image with a logistic regression (LogReg) score below 0.5 is classified as a P5175 cell. (B) Receiver Operator Curve (ROC) of the LogReg model demonstrating a correct single-image classification score of 78%. The area under the curve (AUC) of the ROC equaled 0.847. (C) Absolute value of the regression coefficient associated with the descriptors. This is a measure of the relative importance of each descriptor in the trained LogReg machine learning model. (D) Effect of Recursive Feature Elimination (RFE) on the classification performance of the LogReg model. Performance dropped when less than 21 descriptors are used (dotted line). (E) Background-corrected image of a TMRM-stained CT5120 fibroblast (top panel) and magnification of a region-of-interest (white box). The lower three panels show a “mitogram” depicting all mitochondrial objects in the top panel sorted according to the numerical value (indicated for typical objects) of the “Perimeter ratio”, “Margination” and “Area on Box” descriptor.

machine learning model. To express the classification performance of the trained machine learning model, a score was used that reflects the probability that a given image in the test set is correctly classified as a CT5120 or P5175 cell. The performance of five popular machine learning algorithms was compared (Supplementary Information): Logistic Regression (LogReg), Linear Discriminant Analysis (LDA), Partial Least Squares Discriminant Analysis (PLSDA), Quadratic Determinant Analysis (QDA) and Support Vector Machines (SVMs). Best results were obtained with the LogReg and SVM model using the median per cell data-set (Supplementary Table S2). The LogReg model was used in the remainder of the study since it is computationally simpler than SVM and easier to interpret. The LogReg model condenses the total information contained in the 31 descriptors into a single parameter: the LogReg score. The latter represents the probability that a cell is classified as being CT5120 (LogReg value close to 1) or P5175 (LogReg value close to 0). The LogReg score is displayed against the first principal component (PC1), which captures the maximal variance in the overall data (Fig. 2A). This demonstrates that CT5120 and P5175 cells separate along the LogReg score, whereas no separation is observed along the PC1 axis, meaning that: (i) the difference in morpho-functional phenotype between CT5120 and P5175 cells is captured by the analysis (LogReg) and, (ii) non-supervised machine learning analysis is not appropriate to detect this difference (PC1). The LogReg model is designed in such a way that it takes a binary decision (i.e. any image scoring below 0.5 corresponds to P5175 cells whereas a score above 0.5 corresponds to CT5120 cells), leading to a correct classification rate (per individual image) of 78% on a blind test set. The corresponding area under the curve (AUC) of the receiver operator characteristic (ROC) curve (Fig. 2B), which has as maximal value of 1, equaled 0.85. This value reflects that the ROC-curve is above the line y = x, meaning that the LogReg model has predictive power. These results demonstrate that the two cell lines display observable differences and that the LogReg model is able to classify CT5120 and P5175 cells with 78% accuracy (based upon a single image). To further validate this model, two additional control cell lines from other individuals were analyzed (CT5118: n = 79 images, o = 16860 objects, N = 3 biological replicates; CT5119: n = 109, o = 31153, N = 4). The trained LogReg model also correctly classified these cell lines meaning that the mitochondrial morpho-functional fingerprint of all three control cell lines was classified as identical. Next, we estimated the relative importance of each descriptor in the model, as represented by the absolute value of the LogReg coefficients (Fig. 2C), using a backward elimination similar to the Recursive Feature Elimination (RFE) strategy. In brief, the least informative descriptor (i.e. the one with the lowest LogReg coefficient) was removed from the
The in vitro and cellular antioxidant properties of Trolox and newly developed Trolox variants. As a measure of in vitro antioxidant capacity, we determined the oxygen radical absorbance capacity (ORAC) value. This parameter decreased in the order: KH003 > Trolox > KH001 > KH002 > KH004. Analysis in LS-patient cells revealed that all antioxidants (500 μM, 72 h) reduced the increased levels of CM-H2DCF oxidizing ROS (Fig. 4A). Trolox and KH004 did not affect the elevated levels of HEt (hydroethidium)-oxidizing ROS in LS-patient cells (Fig. 4A). These ROS levels were increased in KH001-treated cells and reduced by KH002 and KH003. None of the compound treatments affected cell shape or adherence (Fig. 4B; typical example of KH003-treated LS-patient cells).

Effect of Trolox and newly developed Trolox variants on the maximal activity of complex I, complex IV and citrate synthase. We previously demonstrated in CT5120 cells that Trolox treatment (500 μM, 72 h) increases the maximal biochemical activity (Vmax) of key mitochondrial enzymes like CI, complex IV (CIV) and citrate synthase (CS)12. Here we observed that Trolox and its four variants displayed similar stimulatory effects in CT5120 cells (Fig 4A). However in case of LS-patient cells, only Trolox and KH003 displayed similar stimulatory effects in CT5120 cells (Fig 4A). In LS-patient cells, only Trolox and KH003 displayed similar stimulatory effects in CT5120 cells (Fig 4A).

Discussion
The case study presented here illustrates the potential of phenotypic screening assisted by objective multivariate statistical methods. Machine learning techniques are particularly well suited to discriminate between phenotypes that are not easily described by a few parameters13 and were successfully applied previously for phenotypic...
profiling of drug effects. The latter study applied a blind approach combining 11 probes to cover maximally the cellular biology and simultaneously testing complete dose response. This untargeted approach was developed in the context of non-specific drug screening.

For analysis of mitochondrial morphology unsupervised learning was used to define six morphological phenotypes in CHO cells and the effects thereupon of caspase inhibition. Similarly, supervised learning was used to analyze mitochondrial morphology in BEAS-2B cells treated with CI and CIV inhibitors and combined with fuzzy logic methods to investigate the link between mitochondrial morphology, ΔΨ and Bax activation. These two last studies use a supervised strategy for mitochondrial morphology evaluation based on three predefined mitochondrial shapes (tubular, donut-shaped or swollen). This means that training was performed using a set of manually chosen images, which might introduce an undesirable selection bias. Our study differs from the above ones since it does not assume the existence of pre-defined mitochondrial morphology phenotypes. Moreover, as far as we are aware, supervised machine learning has not been previously used to evaluate the effects of small molecule treatment on mitochondrial morphology and ΔΨ in a human disease model (i.e. LS cells).

To provide numerical data for machine learning, microscopy images of cells stained with the ΔΨ-sensitive cation TMRM were manually acquired and automatically quantified using a well-established and extensively validated protocol. For every mitochondrial object in each microscopy image a set of 31 descriptors was calculated ("mitochondrial morpho-functional fingerprints"). Relative to a typical healthy control cell line (CT5120), typical LS patient cells (P5175) displayed alterations in 23 out of 31 descriptors. We evaluated various machine learning models for their ability to correctly classify CT5120 and P5175 cells. Although still supervised, the obtained LogReg model is based upon two objectively defined groups (i.e. cells were derived from a distinct healthy subject and an LS patient). A minimum of 21 descriptors was required to maintain the predictive power of the LogReg model. This means that the changes affecting the morpho-functional fingerprints can only be considered from a multivariate perspective. With respect to their average median value, the first two descriptors were not significantly different between CT5120 and P5175 cells (Supplementary Table S1). This demonstrates that parameters that appear uninformative using univariate statistics can be highly relevant in a multivariate analysis, which also considers the relationship between individual descriptors. Biologically, an increase in Perimeter ratio and Area on box indicates that mitochondrial objects and/or (sub)networks become smaller (Fig. 2E). Margination is an intensity-dependent parameter that reflects the homogeneity of the TMRM fluorescence between the center and edge of an object (Fig. 2E and Supplementary Table S1). This means that mitochondrial objects and/or (sub)networks with a high Margination value display inhomogeneous TMRM fluorescence, compatible with spatial ΔΨ inhomogeneities.

In a proof-of-principle study, the developed machine learning strategy was applied to phenotypically classify the effect of the reference compound Trolox and four newly developed Trolox derivatives (KH001–KH004) in P5175 cells. Trolox is a more water-soluble variant of the widely used antioxidant α-tocopherol. Structurally, Trolox consists of a chroman headgroup and a short side chain (Fig. 3A). Both Trolox and α-tocopherol exert their antioxidant activity by reacting with various ROS and lipid hydroperoxyl (LOO•) radicals via the hydroxyl group at position 6. The resulting tocopheryl radical is resonance-stabilized and can be converted back to α-tocopherol by ascorbate leading to formation of an ascorbyl radical. The latter radical species is regenerated by the glutathione (GSH) system. We here designed Trolox variants in such a way that only its side chain was modified whereas the hydroxyl and the methyl groups of the chroman ring were left unaltered. By doing so, we created Trolox variants with different lipophilicity (and thereby water/lipid solubility, cell partitioning and mitochondriotropic character), potentially affecting antioxidant potency, ROS specificity and biodistribution. Visual inspection of the morpho-functional fingerprints reveals that Trolox and its four derivatives all display mitigating effects (Fig. 3B). However, only KH003-treated cells were classified as CT5120 cells by the trained machine learning model. Predictive cheminformatics analysis revealed that KH003 was chemically most similar to Trolox. In _in vitro_ ORAC measurements pointed at KH003 as a slightly more potent antioxidant towards peroxyl radicals than Trolox. Activity analysis showed that Trolox and KH003 stimulated the Vmax of key mitochondrial enzymes (CI, CIV and CS), whereas KH001, KH002, KH004 were less effective. At the cellular level, Trolox and KH001–KH004 reduced the elevated...
levels of CM-H$_2$DCF oxidizing ROS in various LS patient fibroblasts. In the same cell lines, Trolox and KH004 did not reduce the increased level of H$_2$E oxidizing ROS, whereas KH003 reduced HEt oxidation, and KH001 displayed a pro-oxidant effect. These results demonstrate that modification of the Trolox side chain alters its ROS-scavenging properties. Moreover, they support the cheminformatics analysis and machine learning results by highlighting KH003 as displaying the most favorable properties at the cellular level. In summary, we conclude that supervised machine learning is a powerful method to evaluate small molecule effects on mitochondrial morphology and Δψ. Moreover our results suggest that Trolox-derived antioxidants are promising candidate molecules in therapy development for (ROS-related) mitochondrial disorders.

**Methods**

**Cell lines, culture conditions and enzyme activity analysis.** Primary human skin fibroblasts were obtained from healthy individuals (CT5119, CT5118, CT5120) and various Leigh syndrome (LS) patients (P) with isolated mitochondrial complex I (CI) deficiency. Complexes (Oxidative phosphorylation) were predicted and molecularly characterized at the genetic, biochemical and cellular level$^{[40]}$. Cells were cultured under standardized conditions (see Supplementary Information for details). The activities of CI complex IV (CIV) and citrate synthase (CS) were determined in mitochondria-enriched fractions as previously described$^{[46]}$.

**Live-cell microscopy.** Mitochondrial morphology and membrane potential were analyzed in cells stained with the fluorescent cation TMRM (tetramethylrhodamine methyl ester). Cells were visualized using digital imaging microscopy as described in detail in the Supplementary Information. TMRM was used in non-quenching mode$^{[46]}$. Reactive oxygen species levels were analyzed using the ROS-sensing reporter molecules 5-(and 6-)chloromethyl-2, 7'-dichlorodihydro-fluorescein (CM- H$_2$DCF) and hydroethidium (HEt), as described in the Supplementary Information.

**ORAC assay.** The ORAC assay measures the in vitro peroxyl radical scavenging potential of molecules. The antioxidant Trolox is used as a standard in this assay. For details see the Supplementary Information.

**Synthesis of Trolox derivatives.** Detailed information regarding the chemical synthesis strategy is provided in the patent application (Int. patent appl. no. PCT/ NL2013/050528).

**Chemoinformatics, data handling, image quantification and machine learning.** Physicochemical parameters were predicted and molecularly characterized at the genetic, biochemical and cellular level$^{[40]}$. Cells were cultured under standardized conditions (see Supplementary Information for details). The activities of CI complex IV (CIV) and citrate synthase (CS) were determined in mitochondria-enriched fractions as previously described$^{[46]}$.

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**Author contributions**
L.B., J.S., S.v.E., C.V., M.P., L.M.C.B., J.B., P.W. and W.K. wrote the main manuscript. L.B., J.S., M.P., J.B., L.M.C.B., P.W. and W.K. designed the experiments. L.B., S.v.E., C.V., M.P., J.B. and W.K. analysed the data. L.B., S.v.E., C.V. and W.K. prepared the Figure 1. L.B., W.K. prepared the Figures 2 and 3. L.B., M.P. and W.K. prepared the Figure 4. A.J. and R.R. performed the measurements of maximal biochemical activity of Citrate Synthase and Complexes I and IV. L.B. and W.K. performed the data analysis and the prediction of the physicochemical properties of the compounds. S.v.E., C.V., M.P. and J.B. performed the microscopic and ORAC experiments.

**Additional information**
**Supplementary information** accompanies this paper at http://www.nature.com/scientificreports

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