Huntington’s disease (HD) is a fatal inherited neurodegenerative disorder. HD is caused by polyglutamine expansions in the huntingtin (htt) protein that result in neuronal loss and contribute to HD pathology. The mechanisms of neuronal loss in HD are elusive, and there is no therapy to alleviate HD. To find small molecules that slow neuronal loss in HD, we screened 1,040 biologically active molecules to identify suppressors of cell death in a neuronal cell culture model of HD. We found that inhibitors of mitochondrial function or glycolysis rescued cell death in this cell culture and in an in vivo HD model. These inhibitors prevented cell death by activating prosurvival ERK and AKT signaling but without altering cellular ATP levels. ERK and AKT inhibition through the use of specific chemical inhibitors abrogated the rescue, whereas their activation through the use of growth factors rescued cell death, suggesting that this activation could explain the protective effect of metabolic inhibitors. Both ERK and AKT signaling are disrupted in HD, and activating these pathways is protective in several HD models. Our results reveal a mechanism for activating prosurvival signaling that could be exploited for treating HD and possibly other neurodegenerative disorders.

Inhibitors of metabolism rescue cell death in Huntington’s disease models

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Huntington’s disease (HD) is an inherited, adult onset, progressive neurodegenerative disorder (1). HD is caused by a polyglutamine expansion (>36 glutamine repeats) in the huntingtin protein (htt) that leads to neuronal dysfunction and death (1, 2). The mechanism(s) by which the polyglutamine expansion in htt leads to HD pathology remain elusive. Numerous mechanisms including transcriptional dysregulation, altered intracellular trafficking, sequestration of critical cellular proteins in aggregates, aberrant caspase activation, and altered energy metabolism have been implicated in HD (2).

HD is a fatal disease with no therapy. To identify potential compounds for development as drugs and to use these compounds to gain mechanistic understanding of HD, we used a screening approach to identify small molecule suppressors of cell death in a cell culture model of HD. In this model, rat striatal neurons that were immortalized by expression of a temperature-sensitive large T antigen were engineered to express a mutant N-terminal, 548-aa fragment of human htt with 120 glutamine repeats to generate the N548 mutant cell line (3). Serum deprivation and a change to the nonpermissive temperature (39°C) causes T antigen degradation and N548 mutant cell death (3). Cell death can be used as an indicator of mutant htt toxicity because the cells expressing mutant htt die faster than parental cells (3). By using a previously described high-throughput assay (4), we discovered that metabolic inhibitors rescued cell death in this cell culture model and in two in vivo HD models. These compounds activated ERK and AKT prosurvival signaling. Furthermore, growth-factor-induced activation of ERK and AKT rescued cell death, thus elucidating a novel mechanism of rescue by these metabolic inhibitors.

**Results**

**Mitochondrial Inhibitor Rotenone Rescues Cell Death in N548 Mutant Cells.** We screened a collection of 1,040 biologically active compounds (see Materials and Methods) by using a previously described high-throughput screening assay (4). We identified that rotenone, an inhibitor of complex I of the mitochondrial electron transport chain (ETC) (5), suppressed cell death in N548 mutant htt-expressing cells. Cell death rescued was confirmed by three independent cell viability assays and over a time course of 7 days (Fig. 1 A–C). Cell death rescued was verified in three independent clonal cell lines expressing N548 mutant htt (data not shown). We excluded the possibility that rotenone rescued cell death by decreasing the expression of htt transgene or altering T antigen levels (Fig. 1D). Because mutant htt aggregation is associated with HD (2), we tested the effects of rotenone on htt aggregation. In this model, occasional cells (<1%) showed visible aggregation by immunofluorescence [supporting information (SI) Fig. 6]. The low propensity of mutant htt to form microscopically visible aggregates may be due to the longer (N-terminal, 548-aa) htt fragment used in this model compared with other models because decreased aggregation of larger htt fragments has been reported in cell culture and in mouse models (6, 7). The percentage of cells with aggregates was not enhanced by rotenone, indicating that the effect on visible aggregation is unlikely to play a role in cell death rescued by rotenone.

**Diverse Metabolic Inhibitors Rescue Cell Death.** Because rotenone inhibits mitochondrial complex I of the ETC, we expanded our analysis and tested whether perturbing other aspects of metabolism would prevent cell death. We found that diverse inhibitors of metabolism prevented cell death. The compounds tested, their sites of action (Fig. 1E and SI Fig. 7), and their efficacy are shown in SI Table 1. We found that inhibitors of glycolysis (sodium fluoride), ATP synthetase (oligomycin), and mitochondrial coupling 2,4-dinitrophenol (8–10) all rescued cell death. These results indicated that mitochondrial ETC inhibition per se was not required for the rescue.

**Rotenone Rescues Neuronal Loss in Multiple in Vivo HD Models.** Next, we tested the ability of rotenone to alleviate neuronal loss and degeneration in two invertebrate models of HD. In a Caenorhabditis elegans model, an N-terminal, 171-aa fragment of human htt with 150 glutamines is expressed in ASH sensory neurons of polyglutamine enhancer-1 (pge-1) genetic background animals. The pge-1 background enhances mutant htt toxicity and causes age-dependent ASH neuronal death within 3 days (11). ASH neuronal death is monitored by the loss of GFP.
Histone deacetylase inhibitor that is active in Drosophila rotenone was similar to that by sodium butyrate (Fig. 2), with vehicle-treated controls (Fig. 2). Treatment with rotenone rescued rhabdomere loss compared with control treatments (Fig. 2). Animals with a double mutation in S. discans (Sdm) with DMSO or rotenone (10 μM), and cell viability was analyzed every day for 7 days by the trypan blue dye-exclusion assay. The data are averages ± SD of an experiment excluding this mechanism of rescue (SI Fig. 10).

Alterations in ATP, NADH, and Reactive Oxygen Species Levels Are Not Involved in Rescue. Mitochondrial inhibitors can prevent cell death by altering cellular ATP levels (19, 20). We examined whether this mechanism contributed to suppression of cell death in our model. Mitochondrial inhibitors at concentrations that resulted in maximum rescue of cell death caused no decrease in ATP levels in N548 mutant cells (Fig. 3C). Furthermore, time course experiments failed to show a transient decline in ATP levels in oligomycin-treated cells relative to vehicle (DMSO)-treated cells (Fig. 3D). The decline in ATP levels at later times (48 h) is likely due to decreased viability because a larger decline in ATP levels was noted in DMSO-treated cells relative to oligomycin-treated controls.

The result that cellular ATP levels did not decrease upon treating cells with mitochondrial inhibitors was unexpected. One explanation for this result could be that glycolysis was maintaining cellular ATP levels because both glycolysis and oxidative phosphorylation contribute to cellular ATP (SI Fig. 7). We found that oligomycin dramatically decreased cellular ATP levels when glycolysis was inhibited (cells grown in media without glucose) but not when glycolysis was functional (SI Fig. 9). This result suggests that both glycolysis and mitochondrial oxidative phosphorylation contribute to ATP production in N548 mutant cells and that glycolysis was likely maintaining cellular ATP levels upon mitochondrial inhibition.

Mitochondrial ETC inhibition also causes an accumulation of NADH (21), the principal product of the citric acid cycle, and can alter reactive oxygen species levels (21, 22). Increased NADH levels can prevent regeneration and cell death in certain models (21, 23). We found that neither exogenous NADH nor NAD+ nor a precursor of NADH (nicotinamide) rescued cell death at any of the concentrations tested (SI Fig. 10 and data not shown). Furthermore, mitochondrial uncouplers and glycolysis inhibitors rescued cell death (SI Table 1), despite the fact that they do not cause NADH accumulation (SI Fig. 7). Together, these data suggest that accumulation of NADH is not a relevant mechanism of rescue. Finally, we found no changes in reactive oxygen species levels upon treating cells with mitochondrial inhibitors, thus excluding this mechanism of rescue (SI Fig. 10).

Metabolic Inhibitors Activate Prosurvival ERK and AKT Signaling. Altered cell survival signaling (ERK and AKT) is implicated in HD; components of these pathways are decreased or inhibited in HD (24–26), and their activation is protective in HD models (24, 27). We tested whether these pathways were activated by metabolic inhibitor treatment by measuring activating phosphorylation of specific amino acid residues on AKT and ERK proteins (24, 27). Rotenone treatment caused rapid and sustained acti-
HD flies were treated with vehicle (0.1% DMSO), rotenone, or sodium butyrate (NaB). Sodium butyrate (positive control) increased the number of visible rhabdomeres per ommatidium, suggesting that this activation contributed to the rescue of cell death. We also confirmed activation of these pathways by rotenone in two additional clones of N548 mutant cells (Fig. 4D). This activation was not observed in cells treated with vehicle control (DMSO), confirming that it was not an indirect effect of serum starvation (Fig. 4A–D). Furthermore, the pan-caspase inhibitor Boc-D-fmk (28) suppressed cell death of N548 mutant cells (4) but failed to activate these pathways (Fig. 4E), indicating that ERK and AKT activation was not an indirect effect of cell survival. Similar activation of ERK was observed by using the glycolysis inhibitor sodium fluoride (NaF) and the mitochondrial ATP synthetase inhibitor oligomycin, indicating that this activation was not an off-target effect of rotenone treatment (SI Fig. 11).

**ERK/AKT Activation by Rotenone Is Independent of Mitochondrial Respiration.** Our results indicated that a decrease in total cellular ATP levels was not responsible for activating prosurvival signaling or the protection by rotenone. However, because cellular ATP levels are dependent on both glycolysis and oxidative phosphorylation, it was possible that inhibiting mitochondrial ATP production triggered ERK/AKT activation. To test this possibility, we generated two mitochondrial DNA-deficient (ρ−) N548 mutant cell lines by using established protocols (29). In ρ− cells, mitochondrial DNA-encoded subunits of ETC complexes are not generated, making these cells deficient in ATP production by oxidative phosphorylation (30). Rotenone activated ERK and AKT and also rescued cell death in ρ− cells to a similar extent as that in the parental N548 mutant cells (SI Fig. 12). These results indicate that decreased mitochondrial ATP production upon rotenone treatment is unlikely to cause rescue of cell death or to activate ERK and AKT.

**ERK and AKT Activation Partially Explains Rescue by Metabolic Inhibitors.** If activation of ERK and AKT by rotenone is important for rescue, then inhibiting their activation should abrogate the rescue by mitochondrial inhibitors, and, conversely, activating them should rescue cell death. We found that AKT and ERK inhibition by using specific chemical inhibitors of ERK and AKT abrogated the rescue by rotenone in a dose-dependent manner (SI Fig. 13). Moreover, the IC50 of abrogation of rescue by rotenone by these chemical inhibitors was similar to their reported cellular IC50 for the respective kinases (see Materials and Methods), suggesting that they were acting by their proposed...
mechanisms. Next, we tested whether activating ERK or AKT independently could rescue cell death. We used two growth factors, insulin-like growth factor 1 (IGF-1) and EGF-1 which are known activators of AKT and ERK, respectively (31, 32). As expected, IGF-1 caused a sustained activation of AKT but a transient activation of ERK (Fig. 5 A and D). In contrast, EGF-1 treatment caused rapid (15 min) and sustained (∼10 h) activation of ERK and a transient, but lesser, activation of AKT (Fig. 5 B and E). Cells treated with IGF-1 showed a slight rescue (∼5% increased viability), whereas EGF-1 increased viability by 10% (Fig. 5 C). The rescue correlated with a small decrease in caspase 3 activity (Fig. 5 D and E). However, the rescue by growth factors was less than that by rotenone, which increased viability by 30% and substantially inhibited caspase 3 and 7 activation (Figs. 1B and 5 D and E). Because the rescue by these growth factors individually was less than that by rotenone, we tested whether a combination of IGF-1 and EGF-1 would recapitulate the rescue by rotenone. Together, IGF-1 and EGF-1 increased viability by 15% (Fig. 5 C), suggesting an additive effect that was half as effective as rotenone (Fig. 1B). These results suggest that approximately half of the protective effect of rotenone can be explained by activation of ERK and AKT, with the ERK pathway being the major contributor to the rescue.

**Discussion**

By using a high-throughput screen, we discovered that small molecule inhibitors of metabolism (mitochondrial and glycolytic function) suppressed cell death in a striatal cell culture model of HD. Furthermore, several of these inhibitors were also efficacious in *C. elegans* and *Drosophila* HD models (SI Table 1 and Fig. 2), suggesting that this approach targets a conserved mechanism of HD toxicity. The role of energy metabolism in HD is controversial. Several studies implicate deficient energy production in HD pathophysiology; these data have been reviewed elsewhere (1, 33). However, data exist that are not readily explained by this hypothesis. First, several studies in HD animal models and patients show no changes or even higher metabolic rates and/or mitochondrial physiology; these data have been reviewed elsewhere (1, 33). However, data exist that are not readily explained by this hypothesis. Second, HD transgenic...
mice are more resistant to a mitochondrial complex II inhibitor (3-NP) than WT mice (39). Third, caloric restriction decreases energy metabolism (40, 41) and yet alleviates disease progression in HD mouse models (42). Finally, a recent study has identified multiple metabolic changes in HD patients that suggest a procataplastic phenotype (43). Together with our results, these data indicate that decreased energy metabolism may not be critical for the HD phenotype and that a more complex interplay likely exists among mutant htt, energy metabolism, and disease pathology.

The mechanism by which these metabolic inhibitors rescue cell death does not involve a decrease in cellular ATP levels (Fig. 3C) but involves caspase inhibition and activation of prosurvival signaling (Figs. 3B and 5 C–E). Caspase activation is implicated in HD pathology because caspase inhibition shows therapeutic value in mouse HD models (44). N548 mutant cell death is also caspase dependent because serum deprivation activates caspases and a broad-spectrum caspase inhibitor suppresses cell death (4). Metabolic inhibitors prevented “effector” caspase 3 and 7 activation (Fig. 3B), indicating that caspase inhibition by metabolic inhibitors is relevant to their protective effect. Furthermore, these inhibitors activated ERK and AKT; both of these enhance survival in diverse models (32). Activation of ERK and AKT by the growth factors IGF-1 and EGF-1 (32, 45) rescued cell death (Fig. 5C) and could partially explain the rescue of cell death by metabolic inhibitors in our cell culture model.

We investigated the mechanism by which the metabolic inhibitors activate ERK and AKT. Diverse inhibitors of metabolic function such as electron transport (rottenone), glycolysis (NaF), and ATP synthetase (olignycin) activated ERK and AKT in our cell culture model. These results indicate that the prosurvival response is not limited to ETC inhibition or due to off-target effects of these compounds but is likely triggered by common metabolic effects of these compounds. We excluded potential mechanisms, such as changes in ATP, reactive oxygen species, NADH levels (21, 22), and mitochondrial respiration (Fig. 3 and SI Figs. 10 and 12), in triggering prosurvival signaling or rescuing cell death. Although mitochondria are involved primarily in energy metabolism and apoptosis (46), a role for mitochondria in sensing cellular environment and conveying signals to other cell compartments is emerging (47). Understanding the mode of prosurvival signal activation in this cell culture model could reveal conserved links between metabolism and cell survival signaling. Altered growth factor signaling has been implicated in HD pathology; inhibition of EGF-1 and IGF-1 signaling is reported in HD models and patients (25, 26, 48). Conversely, activation of growth factor signaling protects cells from mutant htt toxicity (24, 27). Although extracellular administration of growth factors can activate these pathways, this approach is not feasible in brain tissue of whole animals. Our study reveals a novel means of activating ERK/AKT by using small molecules that could be exploited for therapeutic benefit in HD and possibly other neurodegenerative disorders. Of the active compounds, NaF and 2,4-dinitrophenol were at the dilutions suggested by the suppliers. Other antibodies used have been previously described (4). The National Institute of Neurological Disorders and Stroke custom collection of 1,040 compounds was from Microsource Discovery, Inc. (Gaylordsville, CT); and U0126 and MEK1 inhibitor (52, 53), AKT inhibitor VIII, and phosphatidylinositol 3-kinase inhibitor LY294002 (54) were from Calbiochem (La Jolla, CA).

**Western Blotting.** Cells were incubated with the compounds or vehicle (0.1% DMSO) in low serum or in 10% serum-containing media. Cells were harvested, and Western blotting was performed as described previously (4).

**Cell Viability Assays.** Cell viability was assayed by using the calcein acetoxymethyl ester assay (Molecular Probes) as previously described (4). Trypan blue cell viability was performed on N548 mutant cells by using an automated trypan blue (0.4%) dye–exclusion assay (Vi-Cell 1.01; Beckman Coulter, Fullerton, CA) (4). At least 200 cells were counted per sample, and the percentage of trypan blue negative (viable) cells was calculated.

**ATP Assay.** Cells were plated at 10^5 cells per 60-mm tissue culture dish and, after various treatments, were lysed in 200 μl of lysis buffer; ATP levels were measured by using a luminescence-based assay as suggested by the manufacturer (catalog no. K254-200; Biovision, Mountain View, CA). ATP controls were included to ensure linearity of assay. The light signal (integration, 12 s) was measured in a luminometer (Lumat LB9501; Berthold, Nashua, NH), and luminescence was normalized to cell number. The normalization based on protein content (protein assay reagent; Bio-Rad, Hercules, CA) was similar to that based on cell number.

**Reactive Oxygen Species Assay.** The assay kit was used according to the instructions of the manufacturer (catalog no. D-399; Molecular Probes). In brief, 10 μM H2DCFDA dye (2’’,7’’-dichlorodihydrofluorescein diacetate) was added to 1,500 cells in a 384-well plate, and fluorescence (excitation, 490 nm; emission, 535 nm) was measured by using a plate reader (VICTOR3; PerkinElmer, Waltham, MA).

**C. elegans Neuronal Survival Assay.** The C. elegans assay has been described previously (4, 55). In brief, L1 petty-1;Htn-Q150 animals were incubated at 15°C in liquid S media with a food source (E. coli, OP50 strain) containing either a compound or DMSO (1.5%). GFP fluorescence in bilateral ASH sensory neurons was examined by using a fluorescence microscope (excitation, 485 nm; emission, 535 nm). One hundred neurons were scored in >50 animals. For the starvation assay, L1-arrested animals were obtained by continued absence of food because C. elegans larvae do not initiate growth under starvation (55). These experiments were conducted at 25°C.

**Drug Testing in Drosophila.** The Drosophila HD strain (8534) was obtained from the Bloomington Drosophila stock center and has been described previously (14). Animals were maintained on standard fly food at room temperature. Drugs were mixed in the food every 2 days, and flies were fed on the food for the duration of the experiment at 25°C. The rhabdomere number was assessed by the pseudopupil technique (14). Equal numbers of males and females were included in the control (DMSO) and drug-treated samples. Eight to 10 animals were scored for each treatment, and 35 ommatidia were scored per animal. All scoring was performed in a blinded manner, and significance was calculated by Student’s t test.
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**Electron transport chain**

36 ATP/glucose molecule

**Sodium fluoride**

**Glucose**

**glycolysis**

2 ATP/glucose molecule

**Pyruvate**

**acetylCoA**

**TCA**

**CO₂**

**O₂**

**H₂O**

**NADH**

**2 ATP/glucose molecule**

**INTERMEMBRANE SPACE**

**DNP**

**NAD**

**2e⁻**

**H⁺**

**FADH₂**

**O₂**

**H₂O**

**2e⁻**

**H⁺**

**ADP**

**ATP**

**Oligomycin**

**Rotenone**

**MATRIX**
A

\( \rho^0 \)

\#1  \#2  P  P

- cox IV subunit I
- complex II 70kd subunit
- Tubulin

B

\( \rho^0 \)

\#1  \#2  \#1  \#2

- p ERK
- ERK
- p AKT
- AKT

C

Cell viability (%)

\[ \begin{array}{c|cc|cc}
 & DMSO & Rot & DMSO & Rot \\
\hline
\rho^0 \#1 & 70 & 90 & 50 & 60 \\
\rho^0 \#2 & 60 & 80 & 40 & 50 \\
\end{array} \]