Clinically Approved Heterocyclics Act on a Mitochondrial Target and Reduce Stroke-induced Pathology

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Abstract

Substantial evidence indicates that mitochondria are a major checkpoint in several pathways leading to neuronal cell death, but discerning critical propagation stages from downstream consequences has been difficult. The mitochondrial permeability transition (mPT) may be critical in stroke-related injury. To address this hypothesis, identify potential therapeutics, and screen for new uses for established drugs with known toxicity, 1,040 FDA-approved drugs and other bioactive compounds were tested as potential mPT inhibitors. We report the identification of 28 structurally related drugs, including tricyclic antidepressants and antipsychotics, capable of delaying the mPT. Clinically achievable doses of one drug in this general structural class that inhibits mPT, promethazine, were protective in both in vitro and mouse models of stroke. Specifically, promethazine protected primary neuronal cultures subjected to oxygen-glucose deprivation and reduced infarct size and neurological impairment in mice subjected to middle cerebral artery occlusion/reperfusion. These results, in conjunction with new insights provided to older studies, (a) suggest a class of safe, tolerable drugs for stroke and neurodegeneration; (b) provide new tools for understanding mitochondrial roles in neuronal cell death; (c) demonstrate the clinical/experimental value of screening collections of bioactive compounds enriched in clinically available agents; and (d) provide discovery-based evidence that mPT is an essential, causative event in stroke-related injury.

Key words: caspases • cell death • apoptosis • antidepressants • antipsychotics

Introduction

Evidence from multiple systems has established mitochondria as a critical checkpoint in cell death (1–9). In particular, both caspase-dependent and caspase-independent cell death cascades in the central nervous system (CNS) appear to require mitochondrial release of protein factors such as cytochrome c, AIF, and SMAC/Diablo (10–14). The basic mechanisms underlying the release of these factors remains unclear (15, 16). Induction of the mitochondrial permeability transition (mPT) has been one proposed mechanism underlying this release.

Induction of an mPT has been linked to cytotoxicity after pathological insults such as viral-induced cytotoxicity (17, 18), cardiac ischemia-reperfusion injury (8, 19), stroke and excitotoxicity (6, 7, 20–22), trauma (23), and hypoglycemia (24). mPT, which has been defined primarily using mitochondria isolated from heart and liver, is the opening of...
pores in the inner mitochondrial membrane, resulting in free diffusion of solutes <1.5 kD, destroying the proton gradient, generating reactive oxygen species, and releasing accumulated Ca$^{2+}$ (25–28) as well as cytochrome c, AIF, and SMAC/DIABLO (12, 29–32). These findings link mPT to the release of the direct mediators of downstream caspase-dependent and -independent cell death pathway. A comparison of observations in isolated liver and brain mitochondria suggest that “mPT-like” events do occur in brain mitochondria, even if the characteristics of the brain mPT differ in detail from that which occurs, for example, in liver mitochondria (6, 20, 24, 29).

The specific case of cell death resulting from stroke is associated with pathophysiological changes/conditions that, acting in concert, are predicted, largely on the basis of experiments in isolated liver mitochondria, to favor induction of the mPT. These factors include high Ca$^{2+}$, increases in reactive species, free inorganic phosphate, and activation of upstream protein factors, such as bid and bax (33). Most (20, 29, 34–37), but notably not all (5, 38), studies in cultured cells isolated CNS mitochondria, and experimental models of acute neurological injury (e.g., ischemia/reperfusion) are consistent with the existence of an event analogous with the mPT in the CNS.

However, it remains unclear whether mPT is on the causative pathway of cell death, or whether it is a downstream effect related to overall cellular collapse, which includes, for example, oxidative damage to components of the oxidative phosphorylation system (39, 40). Although neuroprotection mediated by cyclosporine A (CsA) was initially cited as evidence for causative involvement of mPT in ischemic injury (20, 21), this is now appreciated to be problematic as CsA also affects calcineurin, the blockade of which has been shown to be neuroprotective (41). Similar “lack of specificity” arguments hold for minocycline and tauroursodeoxycholic acid. Minocycline is a second generation tetracycline antibiotic known to be protective in models of stroke (42, 43), spinal cord injury (44, 45), and neonatal hypoxia-reperfusion injury (46). Although our recent work links minocycline to prevention of mPT-mediated release of mitochondrially sequestered protein factors that facilitate both caspase-dependent and -independent cell death pathways (29, 47), other actions of minocycline have been identified (42, 43), and the use of minocycline to build a case for mPT involvement awaits a more mechanistic analysis of the actions of minocycline. Tauroursodeoxycholic acid is an endogenous bile acid that protects against stroke, but it is known to modulate activities in three major pathways involved in ischemic damage, including mPT, activity of bel-2 family members, and signal transduction pathways (48).

Likewise, support for an obligate role for tbid activation in ischemia-reperfusion injury now exists (49), but evidence exists that tbid facilitates both mPT induction (33) and mPT-independent release of mitochondrially sequestered pro-apoptogenic factors (16, 50). Similar to the clear evidence for an obligate role of tbid activation, there is also evidence of the required Ca$^{2+}$ deregulation, but there is at least one line of experimentation using cultured cerebellar granule cells that argues against an early role for mPT induction.

Studies of excitotoxic injury in these cells often point to the critical event being a delayed, mitochondrionally mediated deregulation of cytoplasmic calcium as the critical failure event at the level of individual cells (5). The mechanisms that follow kainite exposure seem to differ from those that follow NMDA exposure (51) and neither seems to involve mPT as a critical decision point check (51, 52). It is worth noting that these studies are conceptually difficult to extend in vivo as these cells are comparatively resistant to ischemia.

The best direct test of the hypothesis that mPT lies on the causative pathway of clinically relevant cell death comes from the studies of N-Met-Val-CysA, a nonimmunosuppressive analogue of CsA reputed not to interact with calcineurin. This compound reduces infarct size in a rat model of transient focal ischemia (6). This data is strengthened somewhat by evidence that the effects of CsA on protection against infarction may extend temporally in time from the effects of the calcineurin inhibitor FK506 (53). However, the universal acceptance of mPT involvement in stroke remains limited, in part because of the reliance on data from a single drug (54), and the limited availability and characterization of its analogue. Furthermore, CsA is not viewed as a long-term medical option, as the blood brain must be opened (e.g., by needle puncture) for any therapeutic efficacy (53).

Thus, there is a need to show that other characterized agents can modulate mPT induction and protect against cerebral infarction, both to answer this central mechanistic question in the pathogenesis of stroke-related neuropathology and to help reduce its clinical effects. The work presented here addresses these issues.

Given the potential clinical impact of drugs that inhibit mPT, we screened a collection of 1,040 bioactive compounds for their ability to delay mPT induction in isolated rat liver mitochondria. This collection was primarily composed of clinically approved drugs that could readily be moved to late-stage preclinical trials and then into clinical trials (55, 56). The remainder of the collection largely consisted of some analogues of these drugs, and other known bioactive agents (including natural products, toxins, and controlled substances). Thus, in contrast with large, high, and ultra-high throughput screens of combinatorial libraries, this screen emphasized a much smaller compound set focused on known, bioactive compounds, most of which were FDA approved and, thus, are more likely to be minimally toxic and potentially rapidly available for clinical use. Screening of this collection was blind and was designed to identify compounds that act independently of antioxidant activity or ability to chelate Ca$^{2+}$ approaches that have been taken previously by others.

The data obtained (a) provide discovery-based validation of the mPT as lying on the causative pathway of stroke-related neurologic injury, (b) identify potential therapeutics that are already FDA approved, and (c) demonstrate the potential utility of coupling mechanism-based screening with libraries enriched in well-characterized, clinically available agents.
Materials and Methods

Chemicals. The drug collection screened was the National Institute of Neurological Disorders and Stroke Custom Collection from MicroSource Discovery Systems, Inc. Ultra-pure sucrose was obtained from ICN Biomedicals. All other compounds, of the highest purity available, were obtained from Sigma-Aldrich. All substrates used in mitochondrial respiration experiments were dissolved in respiration buffer (see Fig. 2) and brought to neutral pH.

Mitochondrial Isolation. Liver mitochondria were isolated from ~4–mo-old male Fischer 344 × Brown Norway F1 rats by differential centrifugation using sucrose-based buffers (29). Animal protocols were approved by the Institutional Animal Care and Use Committees of Weill Medical College of Cornell University.

Library Screening and Dose Response Analysis. Assays were run in 265 mM sucrose, 2.6 mM Hepes, pH 7.35, and 2.1 mM K-PO₄, pH 7.35. Incubations included 10 mM α-ketoglutarate, 5 mM glutamate/malate, or 5 mM succinate and CaCl₂ (10 μM for α-ketoglutarate and 20 μM for glutamate/malate and succinate) to energize mitochondria (57). Drugs were screened at 10 μM. Final mitochondrial concentration was 0.75 mg protein/ml. Swelling was monitored by following changes in absorbance at 540 nm and 660 μM (A₅₄₀/A₆₆₀) for 1.5 h using a SpectraMax 250 Plate Reader (Molecular Devices; reference 57). For secondary dose response analysis, 2 out of 27 plates and ~3% of remaining data were excluded as technical outliers by visual inspection. Thus, the dose response data on these agents is based on 7–9 replicates (Fig. S1 and Supplemental Materials and Methods, available at http://www.jem.org/cgi/content/full/jem.20032053/DC1).

Simultaneous Measurement of ΔΨ, Oxygen, Swelling, and Ca²⁺ Transport. Simultaneous measurement of ΔΨ, oxygen consumption, swelling, and Ca²⁺ transport was accomplished using a four-channel respiration system (29, 58). Buffers and additions are described in the figure legends. Oxygen uptake, membrane potential, and Ca²⁺ was measured using Clark, TPP⁺, and Ca²⁺-sensitive electrodes, respectively. Absorbance (A₅₄₀) was measured using a diode. All experiments were performed in triplicate. Representative plots are shown in the paper.

Phospholipase A₂ (PLA₂) and Calmodulin Inhibition. The ability of the heterocyclics to inhibit PLA₂ was determined based on a modification of the method of Meshulam et al. (59). The ability of the heterocyclics to inhibit calmodulin function was assessed by examining the effect of these agents on calmodulin-dependent calcineurin activity (Upstate Biotechnology). Detailed methodologies are presented in the supplemental material.

Neurotoxicity Studies on Cultured Primary Cerebral Neurons. Neurotoxicity studies were conducted as described previously (60). Cerebral cortex of mouse embryos at day 15 (E₁₅) were dissected. The right MCA was occluded with a 7-0 nylon filament and treated (i.p. injection with promethazine solution). All animal experiments were conducted in accordance with the Animal Care Committee. Online Supplemental Material. Online supplemental material includes the following. Details on the library screening methods, primary results, and the validation of the methods used (Figs. S1 and S2). Discussion of preliminary structure–activity relationship (SAR) analysis. Graphic comparisons between inhibition of PLA₂ activity and calmodulin activity and protection against mPT activation (Fig. S3). Methods for these supplemental tests and additional literature citations for activity of related agents against spinal cord injury, sepsis and cardiac, liver and kidney damage (e.g., from ischemia–reperfusion). Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20032053/DC1.

Results

Library Screen. Given the potential clinical impact of drugs that inhibit mPT, we screened a library of 1,040 bio-
Available Drugs Protect Mitochondria and Reduce Stroke-induced Pathology

active compounds for their ability to delay mPT induction in isolated rat liver mitochondria (Fig. 1, Fig. S1, and Supplemental Materials and Methods). Of the 23 compounds showing “moderate” protection in the initial screen, 2 were known mPT inhibitors: tamoxifen and trifluoperazine (both “positive controls”).

![Figure 1](image-url)

**Figure 1.** Heterocyclics mediate dose-dependent protection that is robust and not related to therapeutic class. (A) Drug structure, name, and therapeutic class of each heterocyclics-related structure in the initial compound library. Compounds ranked best to worst (from left to right and top to bottom) based on the data in B. As one measure of potential clinical utility, human use approval was examined using a list of U.S. FDA-approved drugs. Drugs currently in clinical use (23 out of 32) have no abbreviations after their name. Where abbreviations appear after the drug names, they refer to the following: not used in humans (NHU; 1 of 32); used in humans, current status and countries of acceptance uncertain (HU; 4 out of 32); and FDA-approved but believed discontinued (DSC; 4 out of 32). (B) Relative areas under the absorbance curve for the 32 heterocyclics studied at 30 μM. Bars are color coded by each drug’s therapeutic class. Minimal protection was observed at concentrations ≤1 μM, and these data were pooled with the no-drug controls to serve as the control dataset. Total control dataset, n = 1207. Area relative to control, 1.14 ± 0.48 (mean ± SD). The mean coefficient of variation (CV) for individual experiments was 32%, and the mean CV within a specific challenge was 15%. Protection observed was statistically significant at P < 0.05 for 28 out of 32 agents (see Table I). Compounds were assayed in triplicate at seven concentrations (30, 10, 3, 1, 0.3, 0.1, and 0.03 μM) against three different challenges (50 μM Ca²⁺/2.5 mM K-PO4, 25 μM Ca²⁺/5 mM K-PO4, 25 μM Ca²⁺/2.5 mM K-PO₄/100 μM tert-butyl hydroperoxide). Loss of absorbance (i.e., induction of PT) was followed as described in Materials and Methods, and areas under the curve were determined. The increased area under the curve is proportional to delay of PT induction. The area under the curve is expressed as a ratio to the area under the control curves on the same plate. For each drug, n = 7–9. Thick horizontal lines show mean and mean ± 1 SD.
Potency Studies of Heterocyclic, Tricyclic, and Phenothiazine-derived (HTPD) mPT Inhibitors. Inspection of the structures of the 23 compounds that showed moderate protection revealed that trifluoperazine and 12 additional compounds were from a specific subclass of heterocycles and their structural analogues, a major class of psychotropic drugs used clinically since the 1950’s. These included tricyclic antidepressants and phenothiazine-derived antipsychotics. The compound library was reexamined to identify potential analogues that had been considered inactive. This was done both to determine if these were initially false negatives and to identify structurally related, but inactive compounds for SAR analysis. Supplemental Materials and Methods. Retrospective analysis showed that 32 compounds in the collection had a common chemical motif that includes or approximates the tricyclic/heterocyclic backbone (with either a six- or seven-member central ring; Fig. 1 A). Each of these 32 compounds was reassayed against three models of mPT induction and ranked according to the resistance it conferred to induction (Fig. 1 B). Challenges were combined to give a single score reflecting overall protection against mPT (Fig. 1 B and see Table I). Of the experimental set of 32 compounds, 28 gave statistically significant protection at concentrations ≤30 μM. Of these 28, 21 (75%) and 12 (43%) were protective at 10 and 3 μM, respectively. Analytical controls for dose and scoring method are provided as supplemental material (Fig. S2, A and B, available at http://www.jem.org/cgi/content/full/jem.20032053/D11).

Searches of the scientific literature and the US FDA-approved drug list indicate that 23 of these 32 agents are in clinical use, and 4 others are approved but no longer in active clinical use. Four others were approved, at least for clinical trials, in at least one country. Only one (methiothepin) appears not to have been used on humans (Fig. 1 A).

HTPD mPT Inhibitors Do Not Alter Basic Mitochondrial Physiology. Mitochondrial physiological parameters were assessed to determine whether these heterocyclics and their structural analogues interfered directly with other mitochondrial functions, and whether such interference might underlie their inhibition of the mPT due to a nonspecific effect on aspects of mitochondrial function needed for the assay to measure mPT induction. Specifically, we tested three major physiological functions of mitochondria as follows: their ability to respire, their ability to retain a membrane potential (ΔΨ), and their ability to take up and retain exogenous calcium. Representative data under conditions favoring mPT induction are shown in Fig. 2. Compounds shown represent some of the structural and functional diversity present in the heterocyclic compound class (flu- fenazine and promethazine [antihistaminics]; methiothepin [serotonin modulator]; and clomipramine [antidepressant]). At 10 μM, these compounds protected against mPT induction (Fig. 2 D) without apparent effects on initial oxygen consumption before mPT induction (Fig. 2 A), Ca2+ transport (Fig. 2 B), or resting or recovered ΔΨ (Fig. 2 C). These results suggest that the protection against PT is not

![Figure 2](https://example.com/image.png)

**Figure 2.** Heterocyclics delay mPT induction in isolated liver mitochondria without impairing mitochondrial physiology. TPP+ was titrated into buffer to enable construction of standard curves (0–300 s). Succinate-energized mitochondria were added to buffer (with 10 μM of drugs, where noted) within 10–20 s of the graph’s origin (300 s). 100 μM Ca2+ was added where noted, and the curves were standardized at this point. (A) Oxygen consumption was measured by Clark electrode. Oxygen concentration in buffer decreases as signal decreases. (B) Ca2+ transport measured by Ca2+-selective electrode. Ca2+ in buffer increases as signal increases. (C) ΔΨ measured using a TPP+ electrode. ΔΨ decreases as signal increases. TPP+ uptake was determined by comparison with standard curve after correction for shift in electrode signal induced by drugs. Mitochondria appear to take up equal levels of TPP+ in the presence and absence of drug and restore TPP+ and Ca2+ levels after recovering from Ca2+ pulse, suggesting that these drugs have minimal or no direct effects on ΔΨ. Changes in ΔΨ consistent with the slight increases in respiration (equivalent to ~5% of maximally uncoupled mitochondria) observed with some drugs is likely below the limit of detection. (D) Swelling (absorbance) monitored by a light emitting diode at A660. Mitochondrial are more swollen when signal decreases. Assay in 300 mM sucrose, 2.5 mM K-PO4, 3 mM Hepes, pH 7.2, and 5 mM succinate. Experiments were stopped after oxygen was consumed.
associated with any overt mitochondrial toxicity. The addition of the high Ca\(^{2+}\) dose induces mPT in the control sample; drug-treated samples are protected against MPT. Indeed, $\Delta \Psi$ is maintained, swelling is prevented, and calcium is sequestered until the oxygen in the chamber has been consumed. The protective effects of these drugs at the level of purified mitochondria reflect a novel cellular site of action for these drugs, and the ability of these compounds to inhibit the mPT suggests that they represent potential inhibitors of some pathways of apoptosis and necrosis.

**HTTPD PT Inhibitors Do Not Mediate Induction by Inhibiting PLA\(_2\) or Calmodulin.** One heterocyclic/phenothiazine, trifluoperazine, has been identified as an mPT inhibitor and has been shown to be protective against ischemia-reperfusion injury in multiple models. Trifluoperazine can inhibit both PLA\(_2\) (67) and calmodulin (68). Both of these activities have been hypothesized to be protective against mPT. In particular, PLA\(_2\) inhibition has received substantial attention, despite some evidence arguing that it is not the mechanism by which trifluoperazine protects against mPT (67). Literature analysis suggested that other tricyclics and phenothiazines might also inhibit PLA\(_2\) or calmodulin activity. Therefore, we assayed the capacity of each tricyclic in the library to inhibit PLA\(_2\) or calmodulin at concentrations from 1–300 μM. The ability to inhibit either PLA\(_2\) or calmodulin did not correlate with the ability to inhibit mPT in the compound set tested (PLA\(_2\); $r^2 = 0.04$ at 100 μM; calmodulin: $r^2 = 0.02$ at 300 μM; see Table I and Fig. S3, available at http://www.jem.org/cgi/content/full/jem.20032053/DC1).

**Physiological Studies of HTTPD PT Inhibitors.** Because of the rate of apoptosis in stroke-mediated pathology, the activity of a heterocyclic compound was next examined in two models of stroke: (a) oxygen-glucose deprivation (OGD) of cultured primary cerebrocortical neurons; and (b) in the mouse, middle cerebral artery (MCA) occlusion (MCAO)/reperfusion.

Promethazine was chosen as the compound representing this class for three reasons. First, of the most potent inhibitors of mPT, promethazine has one of the highest tolerated doses in humans; although trifluoperazine is the strongest mPT inhibitor, it is used in humans at doses 15-fold below those of promethazine. Furthermore, trifluoperazine has limited utility for long-term use in humans because of its toxicity in nonneural tissues. Although irrelevant for the short-term treatment required by stroke, this limitation would present complications for the treatment of chronic neurodegenerative diseases. Methiothepin, which is approximately equivalent to promethazine in potency, has not been used in humans. Second, promethazine is a well-tolerated drug with few adverse side effects, and, as opposed to the heterocyclics, promethazine has comparatively minor neurological side effects. We note that, from our experience with minocycline (29), the in vitro assay can overestimate the levels of drug necessary for protection in cell culture and in vivo. Therefore, concentrations used in the cell assay were chosen based on concentrations of promethazine shown in the literature to be bioactive. Doses chosen for the in vivo studies were also based on the literature. The dose used was one half those shown to be nontoxic in an analysis including cerebellar pathology. Doses were given at 12-h intervals as it is known promethazine has an 8-h half-life in mice. Third, promethazine does not appear to be a strong PLA\(_2\) or calmodulin inhibitor and, thus, promethazine can be used to demonstrate the independence of effects against mPT induction and action on calmodulin or PLA\(_2\).

**Promethazine Protects Cultured Primary Neurons from OGD.** OGD of primary cerebrocortical neurons is a widely used cell culture model of stroke-related pathology. Promethazine inhibited OGD-mediated neuronal death, as determined by reduction of LDH release into the media (~65% less release). Inhibition of neuronal death was observed at submicromolar concentrations of promethazine (Fig. 3, 0.1 μM, $P = 0.05$; 0.5 μM, $P < 0.001$). Caspase-3 activation plays a key role in hypoxia/ischemia-mediated injury. To gain insight into the mechanism of promethazine-mediated neuroprotection, we evaluated whether this compound could inhibit OGD-mediated caspase-3 activation. As expected from an mPT inhibitor, promethazine-treated neurons exposed to OGD showed reduced caspase-3 activation, supporting the hypothesis that the drug interferes with caspase-mediated cell death.

Follow-up studies have indicated that the tricyclic antidepressant nortriptyline, which also inhibits mPT (Fig. 1), is also protective in this OGD model (unpublished data).
Promethazine Reduces Infarct Size and Neurological Impairment after MCAO-Reperfusion. Because promethazine reduced OGD-mediated neuronal death, we evaluated whether it could also ameliorate ischemic damage in vivo, in which activation of the mPT and caspase-3 may play roles. After MCAO, promethazine-treated mice showed a 53% reduction in lesion size compared with saline-treated mice (Fig. 4 A, n = 13, P < 0.005). Both maximal infarct size and variability of the injury were reduced by 65%. Consistent with a reduction in ischemic damage, the neurologic impairment score of promethazine-treated mice was significantly improved 24 h after MCAO (Fig. 4 B, P < 0.05). Cerebral blood flow (measured by a laser-doppler flowmeter) and systemic blood pressure did not differ between the treated and control groups. As expected from studies of other neuroprotectants, the neurologic score was not different at 30 min after ischemia evaluation (Fig. 4 C and references 61, 62). Protection of both neural function and structure was consistent across all individual mice in the analysis (Fig. 4 D, diagonal line emphasizes that all individuals are protected).

Follow-up studies have indicated that the tricyclic antidepressant nortriptyline, which also inhibits mPT (Fig. 1) is also protective in this MCAO model (unpublished data).

Discussion

Defining the Pharmacophore. The heterocyclics and related compounds described here (HTPD) represent a series of structurally related inhibitors that may help better understand the mPT. SAR analysis, based on the final evidence that 28 of 32 HTPD compounds are protective versus 10 out of 1,008 nonheterocyclics suggests that the HTPD backbone’s activity is highly significant within the intact screening set. Our data is most consistent with the active pharmacophore being the basic class structure, rather than a specific subclass as defined either clinically (e.g., antidepres-

sant vs. antipsychotic), biochemically (e.g., ability to inhibit PLA2), or structurally (e.g., noncarbon atom containing rings vs. tricyclic). There is a wide range of potency for the heterocyclics (Fig. 2), but further SAR analysis is currently limited by a lack of inactive compounds having this basic structure in our current library. Of the group studied, four compounds are not true heterocyclics (one noncarbon atom in the “central” ring), suggesting the classical heterocyclic structure itself may not be mandatory. Mefloquine has only two rings, suggesting the three-ring backbone common in this class may be a reflection of the set of compounds tested rather than an obligate structural feature.

mPT in Stroke. These data are in accord with and significantly extend findings of other groups who have provided evidence linking the mPT to excitotoxic and OGD injury in cultured neurons (7, 69, 70) and to damage subsequent to ischemia-reperfusion in intact animals (6). The Matsumoto et al. (6) and Khaspekov et al. (69) papers are of particular note, as they used the N-Met-Val analogue of cyclosporin A, which has been considered not to react with calcineurin. Retrospective literature analysis also indicates that several HTPD drugs had shown some protection against ischemia or against mPT or mPT-like phenomena, but that the existence of a structural class with a common biochemical target (mitochondria) was not previously recognized, and has not been followed up clinically, possibly due to the side effects of the specific agents tested. Specifically, our data allows us, for the first time, to pull together 15 yr of study of these agents to provide the first common linkage of the HTPD compounds that crosses previously recognized clinical, neurochemical target, and structural bounds, and to provide a probable mechanism for their actions in vivo. Thus, as shown in Table 1, the in vivo and cell culture protection mediated by the HTPDs has been replicated already, by at least 22 labs, in at least 10 HTPD drugs in animal models of cerebral ischemia and related
challenges and by at least 8 HTPD drugs in cell culture. We note that agents have been shown to work both before and after ischemia, and have been tested in mice, rats, and gerbils. Some isolated mitochondrial studies on HTPD agents other than trifluoperazine and tamoxifen has also been published. Histamine has been shown to cause swelling of liver mitochondria, and antihistamines to retard it (68), but this finding has not been examined further. Thioridazine has been suggested to inhibit apoptosis and delay mPT induction by acting to reduce mitochondrial oxidative stress (71). Thus, our data help explain and link a series of studies in different research areas by providing a common mechanism to explain the observations in these studies.

Our data also strengthen the case for mPT as a critical event in the causative pathway of stroke-mediated cell death, although some caveats still remain. Protection against mPT

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<td>58</td>
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<td>Amitriptyline</td>
<td>Antidepressant</td>
<td>2.64 ± 0.77</td>
<td>&lt;0.0001</td>
<td>72</td>
<td>80</td>
<td>78, 80</td>
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<tr>
<td>Amoxepine</td>
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<td>2.50 ± 0.60</td>
<td>&lt;0.001</td>
<td>45</td>
<td>29</td>
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<td>Maprotiline</td>
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<td>2.42 ± 0.56</td>
<td>&lt;0.001</td>
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<td>10</td>
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<td>Quinacrine</td>
<td>Other</td>
<td>2.42 ± 1.21</td>
<td>&lt;0.05</td>
<td>121</td>
<td>43</td>
<td>85–87</td>
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<td>Pericazine</td>
<td>Antipsychotic</td>
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<td>&lt;0.05</td>
<td>84</td>
<td>47</td>
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<td>Ethopropazine</td>
<td>Other</td>
<td>2.30 ± 0.69</td>
<td>&lt;0.01</td>
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<tr>
<td>Mianserin</td>
<td>Other</td>
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<td>Cyclobenzaprine</td>
<td>Other</td>
<td>2.12 ± 0.38</td>
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<td>80</td>
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<td>Imipramine</td>
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<td>2.10 ± 0.50</td>
<td>&lt;0.01</td>
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<td>Clozapine</td>
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<td>91, 93</td>
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<td>Antidepressant</td>
<td>1.68 ± 0.56</td>
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<td>79, 81</td>
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<td>93</td>
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<td>Thiothixene</td>
<td>Antipsychotic</td>
<td>1.03 ± 0.21</td>
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<td>71</td>
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<td>Propantheline</td>
<td>Other</td>
<td>0.84 ± 0.39</td>
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<td>87</td>
<td>76</td>
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<tr>
<td>Pirenzepine</td>
<td>Other</td>
<td>0.70 ± 0.30</td>
<td>NA</td>
<td>485</td>
<td>45</td>
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</table>

Drug class as in Fig. 1 B. PT presented as mean ± SD as fold protection over control. Calmodulin (Cal) and phospholipase A₂ (PLA₂) as percent control activity over untreated sample. Cerebral IR lists references for in vivo models are relevant for cerebral ischemia-reperfusion. Cell lists references for cell culture models are relevant for cerebral ischemia-reperfusion.

aThis paper.
bThis paper and unpublished data.
cUnpublished data.
NS, P > 0.05; NA, not applicable, mean < 1.
and in vivo protection against ischemia-reperfusion and related events/challenges were shown to be qualitatively correlated in at least 10 compounds (Table I). All of these compounds displayed activity in at least one study in the range of 3–20 mg/kg, consistent with relatively similar dose dependence in in vitro assays of mPT inhibition. In contrast, protection mediated by these agents does not correlate with their ability to inhibit PLA2 or calmodulin and cannot be attributed to either the clinical or structural class of the mPT inhibitor (Fig. 1, Table I, and Fig. S3). By providing direct evidence consistent with mechanistic involvement of mPT inhibition in the actions of these 10 compounds, removing potential alternative explanations (e.g., PLA2 inhibition, calmodulin inhibition, and neurochemical effects), and providing evidence that mPT inhibitors structurally distinct from CsA are neuroprotective, our data provide direct counter-arguments to each of the major concerns raised previously (see Introduction) against mPT playing a causative role in stroke-mediated damage. Remaining caveats include, for example, (a) the identification of these compounds as mPT inhibitors is still based on work in isolated liver mitochondria, the biochemically defined system, but not the target tissue; (b) there are systems, for example, the calcium overload model (72), that may not involve mPT as a primary mediator; and (c) it is always possible that a previously unrecognized system is the actual target (e.g., the TRMP7 channels recently recognized to contribute to anoxic cell death in some models; reference 73).

Potential Clinical Utility. The results reported here may expand the potential clinical uses of HTPD drugs to protection against stroke, and set the stage for studies further expanding their utility to other disorders. Additional studies are required; however, our data suggest that these agents are potentially usable as long-term prophylactics (to reduce damage from an event that occurs, not to prevent the event) for individuals at risk for strokes and heart attacks (e.g., patients undergoing carotid endarterectomy or who have experienced a cardio–cerebrovascular event). The heterocyclics may also be appropriate for acute management of stroke and heart attack as well as a component of the clinical management of neurodegenerative disease. Further retrospective analysis (see Online Supplemental Material) showed two HTPD compounds at delayed mPT are protective against spinal cord injury (trifluoperazine and chlorpromazine) and three against myocardial and kidney ischemia-reperfusion (trifluoperazine, chlorpromazine, and quinacrine). Chlorpromazine has also been shown to protect against sepsis and liver toxicity/ischemia.

Although mPT involvement in chronic neurodegenerative disorders remains unknown and controversial, the common involvement of mitochondria in cell death pathways suggests that the HTPD drugs might be considered, in future experiments, as potential candidates for long-term use in individuals with neurodegenerative diseases that involve cell death in the nervous system, including amyotrophic lateral sclerosis, and Parkinson’s, Alzheimer’s, and Huntington’s diseases. Applicability in Huntington’s disease is supported by data showing that 5–10 mg/kg promethazine protects against striatal lesions in Lewis rats induced by 3-NP (unpublished data). Applicability in Parkinson’s disease is supported by data showing that promethazine (best at 10 mg/kg) protects against striatal dopamine depletion and neuron loss in the substantia nigra pars compacta induced by MPTP (unpublished data). Both preclinical and clinical testing will be required to determine whether these drugs will be beneficial for these diseases.

These drugs appear to offer major advantages as clinical mitochondrial protective agents. First, 31 out of the 32 compounds tested here are clinically available, and 27 are clinically approved, suggesting the potential for rapid movement into preclinical and/or clinical trials. Some of these drugs have been used for five decades. Second, the active drugs appear from their clinical uses and side effects to cross the blood brain barrier (three possible exceptions being cyclobenzaprine, metloquine, and quinacrine). Third, most are safe for long-term use, and many have only minor side effects (a notable exception being the phenothiazine trifluoperazine). Fourth, they are active in vitro and in vivo at concentrations close to or within those that are achievable in humans. The maximal therapeutic dose of promethazine in humans is 3 mg/kg, and several have LD50 in animals 5–10 times higher than promethazine with favorable safety profiles. Last, the many available classes of these agents with differing side effects and clinical actions suggest that a combination of agents, each one in a subclinical dose, can be used to achieve mPT inhibition without inducing unwanted effects. This work defines a structural family suitable for further pharmacological optimization, complementing the promise of available clinical therapies and novel experimental reagents with a longer range promise of even further optimized agents (e.g., for protection and for probing mitochondrial involvement in cell death in isolated systems, cultured cells, and in vivo). Although further preclinical study is warranted, these findings suggest that these clinically approved drugs will be candidates for human study in stroke and neurodegeneration.

The authors thank J. Moore for assistance with publication graphics.

This work was supported by grants from the Hereditary Disease Foundation (to B.S. Kristal), National Institutes of Health (NIH)/National Institute on Aging (to B.S. Kristal), NIH/National Institute of Neurological Disorders and Stroke (to R.M. Friedlander and A.M. Brown), the Huntington’s Disease Society of America (to R.M. Friedlander), and Burke Medical Research Institute (to B.S. Kristal). No author has direct competing interests in the work presented. A patent application has been filed on aspects of this work by Weill Medical College, and some authors potentially have patent rights though university agreements.

Submitted: 26 November 2003
Accepted: 26 May 2004

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1. Zamzami, N., S.A. Susin, P. Marchetti, T. Hirsch, I. Gomez-


Available Drugs Protect Mitochondria and Reduce Stroke-induced Pathology

Downloaded from on April 5, 2017
Materials and Methods

**PLA2 Inhibition.** The ability of the heterocyclics to inhibit PLA2 was determined based on a modification of the method of Meshulam et al. (60). In brief, the fluorescent PLA2 substrate Bis-BODIPY-glycerophosphocholine (bis-BODIPY FL C11-PC) was incorporated into mitochondrial membranes and fluorescence self-quenching ensues. Separation of the fluorophores on hydrolytic cleavage of one acyl chain by PLA2 results in increased fluorescence. Reaction buffer contained 310 mM sucrose, 3 mM Hepes, pH 7.4, 40 μM CaCl2, 2.5 mM K-PO4, 5 mM succinate, or 5 mM glutamate/malate, 0.1 mM bis-BODIPY FL C11-PC, and 1 mg/ml mitochondria. For determination of the drug’s action on PLA2, mitochondria were treated with the drug’s stocks in concentrations from 0.3 to 300 μM. Kinetics of the fluorescence was monitored for 120 min at excitation, 488 nm, and emission, 530 nm.

**Calmodulin Inhibition.** The ability of the heterocyclics to inhibit calmodulin function was assessed by examining the effect of these agents on calmodulin-dependent calcineurin activity (Upstate Biotechnology). Calcineurin activity was determined by following the absorbance of the product of calcineurin/PP2B-induced hydrolysis of calcineurin-specific substrate p-nitriphenyl phosphate (pNPP) in the presence of appropriate cofactors (Ca, Ni, calmodulin). The specific enzyme activity represents the inverse of the mass of calmodulin required to give half the maximal reaction rate. Determination of calcineurin/PP2B activity included two steps: enzyme activation and the enzymatic reaction. For enzyme activation, calcineurin, 1 μg calmodulin, 0.1 mM CaCl2, 1 mM NiCl2, and 1% BSA were incubated for 15–30 min at 37°C. The enzymatic reaction was started by addition to the reaction mixture of calcineurin substrate (pNPP) and further incubation for 10–30 min at 37°C for color development. Absorbance was monitored at 405–410 nm. For determination of a drug’s action on calcineurin/PP2B activity stocks, the drug (at 0.3–300 μM) was added to the reaction mixture during the enzyme activation stage.

Results

**Library Screening.** Primary screening of the compound library and subsequent follow-up on initial hits identified 37 potentially useful compounds that were mPT inhibitors at 10 μM (Fig. S1). Primary screens were performed using a standard spectrophotometric assay based on mPT-induced swelling of energized liver mitochondria after a Ca2+ challenge (1). Each of the 1,040 compounds were tested against the low calcium challenge three times (once each with glutamate/malate, succinate, or α-ketoglutarate). In the initial screen, drugs were scored as providing “strong” protection (basically no change in absorbance), “moderate” protection (observable delay in loss of absorbance), or no protection. The scoring scheme is pre-

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**Figure S1.** NINDS library screening procedure. (A, top left) View of a screening plate showing raw data. mPT induction was assessed spectrophotometrically essentially as described previously (Materials and Methods and references 1, 9). (top right) Seven compounds of potential interest and six controls are highlighted for demonstration purposes. This view shows change in absorbance from starting point and has advantages for identifying compounds that are relatively weakly protective at the doses tested. (bottom left) Change to absolute absorbance highlights the true versus false positives. (bottom right) Expanded views of one example of each class. Note that all studies are conducted simultaneously at two wavelengths to avoid wavelength specific effects due to compound absorption. (B) Table of overall outline of assay progression for the first five screens in the study. Conditions as noted in the figure legend and text. Each assay was performed once with each substrate. Numbers in “strong”/“moderate” inhibitors reflect the number that showed similar effects against all substrates (first number) or against only one substrate. Screening assays 1, 2, and 3 were plate reader–based assays for loss of spectrophotometric absorption as shown (left). Assay 4 was a respiration assay (Materials and Methods). Assay 5 involved simultaneous analysis of oxygen consumption, membrane potential (ΔΨm), Ca2+ transport, and mPT inhibition. Examples of this assay with additional experimental details are shown in Fig. 2 and described in Materials and Methods.
sented in Fig. S1 A. Compounds protective in at least one substrate were followed as potential inhibitors. Subsequent assays (Fig. S1 B) were performed using varied conditions to overcome protection mediated by compounds acting as chelators (Table S1, line 2) or antioxidants (Table S1, line 3), based on the rationale that we were unlikely to identify novel drug activities in these categories. Remaining drugs were examined to determine whether they provided protection by inhibiting respiration or uncoupling, expected artifacts in our system.

The screen ultimately identified 37 potentially useful compounds that were mPT inhibitors at 10 μM. These remaining compounds (Fig. S1 B, bottom right) are robust inhibitors of mPT in the presence of multiple inducers. These do not act by primarily chelating Ca\(^{2+}\) because the high Ca\(^{2+}\) challenge would overwhelm protection. The protection against high doses of the oxidant tert-butyl-hydroperoxide indicates that these agents are unlikely to be acting primarily either as direct scavenging antioxidants or as specific protectors of the dithiol moiety proposed as an mPT regulator (termed the “S-site”), or the high dose oxidants would readily overwhelm protection. When the blind was broken, it was determined that the 15 strong substrate-independent inhibitors identified in the initial PT screens included only our study’s positive control (cyclosporin A) and 14 compounds with known or determined (not shown) mitochondrial effects that rendered them therapeutically unsuitable or that meant that the results observed were unrelated actions at the level of PT inhibition (e.g., respiratory toxins, uncouplers, other agents with substantial known toxicity at the concentrations used, disinfectants, etc.). In addition to cyclosporin A, these compounds included nigericin, antimycin A, merbromin, bithionol, hexachlorophene, gossypol–acetic acid complex, calcimycin, monesin, lasalocid, salinomycin, alexidine hydrochloride, sparteine sulfate, usnic acid, and rotenone). These agents were not further examined for this paper).

**SAR Analysis.** There is a wide range of potency for the heterocyclics (Fig. S1), but further SAR analysis is currently limited by a lack of inactive compounds having this basic structure in our current library. Overall, the heterocyclics and related compounds described here represent a series of structurally related inhibitors that may help better understand of the mPT. Of the group studied, four compounds are not true heterocyclics (one noncarbon atom in the “central” ring), suggesting the classical heterocyclic structure itself may not be mandatory. Mefloquine has only two rings, suggesting the three-ring backbone common in this class may be a reflection of the set of compounds tested rather than an obligate structural feature.

Preliminary SAR analysis, based on the final evidence that 28 out of 32 heterocyclics and their structural analogues are protective versus 10 out of 1,008 nonheterocyclics suggests that the heterocyclic backbone’s activity is highly significant within the intact screening set.

### Table S1. Protection by Trifluoperazine, Chlorpromazine, and Quinacrine

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<td>Quinacrine</td>
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</table>

Numbers refer to references citing protection in relevant models.

**Figure S2.** Validation for dose response studies. (A) Comparative protection assessed as area under the curve for the 32 drugs at 10 and 30 μM. Shaded box represents no protection and/or enhanced induction. Databased on 23–25 assays at each of these concentrations. (B) Comparative protection assessed as area under the curve and percent initial absorbance remaining when unprotected mitochondria have failed. Databased on 23–25 assays at each of these concentrations.

**Figure S3.** Inhibition of phospholipase A\(_2\) and calmodulin is not related to inhibition of PT induction. (A) PLA\(_2\) activity in presence of 100 μM compound (presented as percent of activity in absence of drug) of 31 of the heterocyclics plotted versus protection against mPT protection (as area under the curve at 30 μM), regression line \(y = 2.9 - 0.006X\), \(r^2 = 0.04\). \(n = 4–6\) for each compound. (B) Calmodulin activity in presence of 300 μM compound (presented as ratio of activity/untreated plate controls) of 28 of the heterocyclics plotted versus protection against mPT protection (as area under the curve at 30 μM), regression line \(y = 3.1 - 0.46X\), \(r^2 = 0.02\). \(n = 3\) for each compound. Loratadine and pirenzepine showed apparent calmodulin activation and were not plotted.