Physiological Phenotype and Vulnerability in Parkinson’s Disease

D. James Surmeier1, Jaime N. Guzman1, Javier Sanchez1, and Paul T. Schumacker2

1Department of Physiology, Northwestern University, Chicago, Illinois 60611
2Pediatrics, Feinberg School of Medicine, Northwestern University, Chicago, Illinois 60611
Correspondence: j-surmeier@northwestern.edu

This review will focus on the principles underlying the hypothesis that neuronal physiological phenotype—how a neuron generates and regulates action potentials—makes a significant contribution to its vulnerability in Parkinson’s disease (PD) and aging. A cornerstone of this hypothesis is that the maintenance of ionic gradients underlying excitability can pose a significant energetic burden for neurons, particularly those that have sustained residence times at depolarized membrane potentials, broad action potentials, prominent Ca2+ entry, and modest intrinsic Ca2+ buffering capacity. This energetic burden is shouldered in neurons primarily by mitochondria, the sites of cellular respiration. Mitochondrial respiration increases the production of damaging superoxide and other reactive oxygen species (ROS) that have widely been postulated to contribute to cellular aging and PD. Many of the genetic mutations and toxins associated with PD compromise mitochondrial function, providing a mechanistic linkage between known risk factors and cellular physiology that could explain the pattern of pathology in PD. Because much of the mitochondrial burden created by this at-risk phenotype is created by Ca2+ entry through L-type voltage-dependent channels for which there are antagonists approved for human use, a neuroprotective strategy to reduce this burden is feasible.

Parkinson’s disease (PD) is a disabling neurodegenerative disorder that is strongly associated with aging, increasing exponentially in incidence above the age of 65 (de Rijk et al. 1997; de Lau et al. 2004). Currently, there is no therapeutic strategy that is proven to slow or stop disease progression. Without a neuroprotective treatment, the incidence of PD is expected to increase dramatically worldwide in the coming decades as life expectancy increases (Dorsey et al. 2007). This will pose an enormous monetary and human burden on society.

The best hope for developing a neuroprotective treatment lies in a better understanding of disease pathogenesis. In the last two decades there has been a great deal of progress in identifying genetic risk factors for PD (Cookson 2010; Cookson and Bandmann 2010; Wider et al. 2010). There is the reasonable belief that identifying the consequences of these mutations will lead to new therapeutic strategies, perhaps ones specifically designed for a particular mutation. However, the vast majority of PD cases are not associated with any known genetic
mutations. Genome-wide association studies (GWAS) promise to put disease risk in these cases on a firmer genetic footing (Gubitz and Gwinn 2009). The working hypothesis of these studies is that the proteins linked to risk-increasing polymorphisms will begin to segregate into well-defined subcellular systems, providing insight into pathogenesis.

Although potentially providing important information about pathogenic mechanisms, these studies will not explain the distribution of neuron loss or pathology in PD. This pattern will be determined by epigenetic factors. Unfortunately, this relationship is not likely to be a simple one, as the expression pattern genes linked to PD do not predict the pattern of pathology. Hence, it must be that other phenotypic factors govern the functional consequences of PD-linked mutations and polymorphisms. Neurons and the networks they form can be viewed as very specialized systems that create the context in which PD-linked mutations manifest themselves. Understanding why neural systems are vulnerable to these mutations could provide a clear path to therapeutic intervention.

This review focuses on one distinctive feature of neural systems, excitability. The energetic demands that attend the maintenance of electrochemical ionic gradients necessary for excitability could be a key determinant of vulnerability. Although plausible, this in and of itself would not point to a therapy, as there is no obvious way to change this characteristic without broadly compromising brain function. But the pathology in PD is not found in all neurons. The most widely accepted pathological marker in PD is the Lewy body (LB) or Lewy neurite (LN), a proteinaceous intracellular deposition (Braak et al. 2004). If LB/LNs are used as a PD marker, it is clear that the pathology is not randomly distributed and far from ubiquitous. In fact, the percentage of brain neurons affected through the mid-stages of the disease appears to be far less than 1%. Moreover, this fraction of LB/LN-containing neurons is not randomly distributed in the brain; rather, these neurons reside in reasonably well-defined nuclei with different functional roles, suggesting that physiological phenotype confers risk. This is an important clue about the mechanisms that bring out the pathological consequences of factors that affect most if not all neurons (i.e., age, mutations in widely expressed genes, and toxins).

**THE NEURONAL PHENOTYPE**

PD is a disease of neurons, not of the liver, kidney, or heart. An implication of this fact is that one or more of the features distinguishing neurons from these other cell types must contribute in a seminal way to pathogenesis. What then distinguishes neurons? A cardinal feature of neurons that separates them from nearly all other cell types is excitability. Neurons use a steep electrochemical gradient across their plasma membrane to perform computations on incoming chemical signals from other neurons and to pass the outcome of this computation to other neurons. Chemical signaling between neurons is for the most part accomplished at specialized junctions between neurons called synapses where a presynaptic transmitter release site is paired with a postsynaptic site at which receptors for the transmitter are aggregated. Effective synaptic transmission between neurons depends on the maintenance of electrochemical gradients for Na\(^+\), K\(^+\), Ca\(^{2+}\), and Cl\(^-\) across the plasma membrane. By gating the flow of one or more of these ions, ionotropic receptors generate electrical signals that are integrated by the postsynaptic neuron. Consider the principal excitatory transmitter in the brain, glutamate. Excitatory synaptic transmission mediated by ionotropic glutamate receptors depends on the electrochemical gradient for both Na\(^+\) and Ca\(^{2+}\). Most AMPA receptors (AMPARs) are selectively permeable to Na\(^+\) ions. When 2-amino-3-(5-methyl-3-oxo-1,2-oxazol-4-yl)propionic acid (AMPA) receptors are activated, they open a pore that allows Na\(^+\) ions to flow into the electronegative cytoplasm, depolarizing the membrane.

N-Methyl-D-aspartate (NMDA) receptors (NMDARs) are permeable to both Na\(^+\) and Ca\(^{2+}\) ions, allowing them to not only depolarize the membrane, but to initiate intracellular signaling cascades dependent on Ca\(^{2+}\). An important feature that distinguishes NMDARs is that they require membrane depolarization (in
addition to ligand binding) to conduct (depolarization dislodges Mg$^{2+}$ ions from the pore). In most neurons, the resting membrane potential is more negative than the voltage at which Mg$^{2+}$ dislodges, effectively reducing the opening of NMDARs during routine excitatory synaptic events.

**SPIKING POSES A METABOLIC BURDEN**

The outcome of the computation performed by a neuron in response to synaptic events is coded in a pattern of regenerative events called action potentials or spikes. Spikes are transient regenerative events that rely on the transmembrane electrochemical gradients for Na$^+$, K$^+$, and Ca$^{2+}$ ions (Fig. 1). In most neurons, the spike is triggered by the opening of voltage-dependent membrane pores or channels that are selectively permeable to Na$^+$ ions, allowing positively charged Na$^+$ ions to move from the extracellular space into the cytosol. This redistribution of charge pushes the transmembrane potential from relatively negative membrane potentials to near 0 mV. This depolarization causes voltage-dependent channels that are selectively permeable to K$^+$ ions to open, resulting in the movement of positively charged K$^+$ ions from the cytosol into the extracellular space. This movement is then counteracted by the activity of pumps and transporters, such as Na/K ATPase, which requires ATP as the energy source to maintain these gradients.

**Figure 1.** Schematic representation of the plasma proteins that regulate ionic gradients following neurotransmitter release or during the time course of an action potential spike. Synaptic input and spiking dissipate ionic gradients maintained by pumps and exchangers creating a metabolic burden. Illustrated in the model are the classic influx and efflux of ions favored by their electrochemical gradients. Particularly, the electrochemical gradient for Ca$^{2+}$ ($\approx +128$ mV under physiological conditions) favors a strong influx of Ca$^{2+}$ ions through NMDA receptors (NR) following glutamate release or activation of Ca$\delta$ channels during pacemaking spiking activity. The tight regulation of intracellular Ca$^{2+}$ is routed into the endoplasmic reticulum (ER) via sarco(endo)plasmic reticulum Ca$^{2+}$-ATPase (SERCA) pumps. However, Ca$^{2+}$ influx into the ER can trigger the process of Ca$^{2+}$-induced Ca$^{2+}$ release (CICR), displayed here as Ca$^{2+}$ released through the IP3R/RyR pathway. Another venue of regulation of Ca$^{2+}$ depends on whether the cell expresses Ca$^{2+}$-binding proteins (CBP), which help buffer intracellular Ca$^{2+}$. Whether CBP is present or not, Ca$^{2+}$ can also be extruded out of the cell via plasma membrane Ca$^{2+}$-ATPase (PMCA) or Na$^+$/Ca$^{2+}$ exchanger (NCX) proteins. SERCA, PMCA, NCX, and the Na/K ATPase all require ATP as the energy source to maintain these gradients, and this ATP consumption can have a strong impact on the bioenergetics of the cell to maintain normal levels of intracellular Ca$^{2+}$.
ions in the opposite direction: from the cytosol to the extracellular space, reestablishing the potential gradient. This sequence of events requires that the concentration of Na\(^+\) ions be low in the cytosol, but the concentration of K\(^+\) ions high.

In most neurons, spikes are initiated in a specialized region at the junction between the cell body and the axon called the axon initial segment (AIS). The AIS is specialized in several respects, most notably in the concentration of voltage-dependent Na\(^+\) channels. Spikes initiated in the AIS propagate both down the axon to synaptic terminals (where they initiate transmitter release) and into the somatodendritic region, creating a spatially distributed dissipation of the transmembrane electrochemical Na\(^+\) and K\(^+\) gradients.

Another cation that crosses the plasma membrane during spikes is the Ca\(^{2+}\) ion. In most neurons, voltage-dependent Ca\(^{2+}\) channels are opened only by strong depolarization during the spike. With repolarization of the membrane potential, these channels slowly close, creating a period during which the driving force for influx of Ca\(^{2+}\) is large and the conductance remains high. This creates a "double hump" in the Ca\(^{2+}\) influx during a spike and makes the total Ca\(^{2+}\) influx during a spike very sensitive to spike duration (Fig. 2). Ca\(^{2+}\) entering during the spike serves a number of ends (Clapham 2007). One is activation of K\(^+\) channels that help repolarize the membrane potential and impede the generation of another spike for some period of time (refractory period). This interaction takes place in a spatially restricted zone close to the plasma membrane. Neurons that need to spike at high frequencies typically restrict Ca\(^{2+}\) entry by keeping spikes very brief (<1 ms); Ca\(^{2+}\) entry during the spike is exclusively dependent on the duration of the spike itself, which varies considerably from one cell type to the next (Fig. 2). To help manage Ca\(^{2+}\), neurons often express fixed Ca\(^{2+}\) buffering proteins (Augustine et al. 2003; Mattson 2007), such as parvalbumin, in addition to the Ca\(^{2+}\) signaling proteins (Clapham 2007).

Exchangers and pumps are responsible for maintaining the electrochemical gradients for

![Figure 2. Spike width shapes Ca\(^{2+}\) entry. (A) Spikes recorded from a substantia nigra pars reticulata (SNr) GABAergic neuron and a substantia nigra pars compacta (SNc) dopaminergic neuron (DA). SNc DA neurons have wider spikes compared to SNr GABAergic neurons. (B) In the top panel, simulated spikes of varying width using the computer program Neuron are shown. In the bottom panel, Ca\(^{2+}\) currents were generated in the model in response to the simulated spikes. Simulated Ca\(^{2+}\) currents were evoked by opening of Ca\(_{\text{v1.3}}\) and Ca\(_{\text{v2.1}}\) model channels in Neuron. The color-coding of the simulated spikes and respective Ca\(^{2+}\) currents show that the amount of Ca\(^{2+}\) charges that enter the cell is higher in simulated spikes with largest width. This model supports our hypothesis that SNc DA neurons that have large spike width will allow more Ca\(^{2+}\) entry compared to an SNr GABAergic neuron, which then can have a strong impact on the undergoing Ca\(^{2+}\) signaling, handling, and metabolic demands for neuronal populations that can differ based on spike waveform.](http://perspectivesinmedicine.cshlp.org/4/4/4/C2A009290)
Na\(^+\), K\(^+\), Ca\(^{2+}\), and Cl\(^-\). These transmembrane proteins fall into two broad categories. The first are pumps that rely on adenosine triphosphate (ATP) to drive the movement of ions. Pumps that fall into this category include the Na\(^+\)/K\(^+\) ATPase, the plasma membrane Ca\(^{2+}\) ATPase (PMCA), and the smooth endoplasmic reticulum Ca\(^{2+}\) ATPase (SERCA). The other category uses the energy stored in an existing electrochemical gradient to move ions. A good example of this type of protein is the Na\(^+\)/Ca\(^{2+}\) exchanger (NCX), which under physiological conditions uses the Na\(^+\) gradient to move Ca\(^{2+}\) ions out of the cytosol. Another exchanger of this type is the Na\(^+\)/Ca\(^{2+}\)-K\(^+\) exchanger (NCKX). Together, this combination of pumps and exchangers maintain the transmembrane ionic gradient for cations (Fig. 1).

Although the molecular events coupling ion movement to ATP hydrolysis are still not fully understood, the thermodynamics of ion movement are worth considering because they establish lower limits on the cost of pumping. For Na\(^+\) and K\(^+\) ions, the concentration differences maintained across the plasma membrane are similar, being 10- to 30-fold. In contrast, the concentration difference for the Ca\(^{2+}\) concentration is \(\sim 20,000\)-fold, being 2 mM in the extracellular space and around 100 nM in the intracellular space. Because the free energy change needed to move an ion from one compartment to another depends on the logarithm of the concentration ratio between the two compartments, Ca\(^{2+}\) should be roughly eight times more energetically expensive than a Na\(^+\) ion (about four times greater on a per charge basis). The Na,K-ATPase extrudes three Na\(^+\) ions and takes up two K\(^+\) per ATP molecule consumed. The PMCA pumps one Ca\(^{2+}\) molecule out for each ATP consumed. So overall, it takes about one ATP for each Ca\(^{2+}\) ion, and one ATP for every three Na\(^+\) ions. By this rough calculation, although Ca\(^{2+}\) is more expensive than Na\(^+\), it is only about 1.5–2 times more expensive per charge than Na\(^+\). Whether the bioenergetic cost of normal Ca\(^{2+}\) influx represents a significant fraction of the metabolic capacity of neurons, remains to be seen.

Ca\(^{2+}\) that is not pumped back out of the neuron rapidly is sequestered in intracellular organelles. The most important of these for Ca\(^{2+}\) homeostasis is the endoplasmic reticulum (ER) (Berridge 2002). The ER forms a continuous, intracellular network, allowing it to regulate both local and global Ca\(^{2+}\) signals. As in other neurons, the ER network in SNc DA neurons extends throughout the somatodendritic tree (Schwyn and Fox 1974; Mogami et al. 1997; Park et al. 2000; Choi et al. 2006). High-affinity ATP-dependent transporters move Ca\(^{2+}\) from the cytoplasm into the ER lumen. The absence of high-affinity, anchored intraluminal Ca\(^{2+}\) buffers and the physical continuity of the lumen within the cell (Mogami et al. 1997; Park et al. 2000) allows the ER to rapidly (~30 μm/s) redistribute Ca\(^{2+}\) between intracellular compartments. Ca\(^{2+}\) sequestered in the ER is released at sites where it can be pumped back across the plasma membrane or where it can be used to modulate cellular function (Rose and Konnerth 2001; Cui et al. 2004; Verkhovsky 2005; Bardo et al. 2006; Park et al. 2008).

Mitochondria partner with the ER in Ca\(^{2+}\) homeostasis. Mitochondrial motility is controlled in such a way as to bring them close to sites of Ca\(^{2+}\) release (Boldogh and Pon 2007; Chen and Chang 2009). Mitochondria also are commonly found physically coupled to the ER by mitochondria-associated membrane (MAM) (Pizzo and Pozzan 2007; Hayashi et al. 2009). These regions of juxtaposition create multifunctional signaling microdomains (Csordas et al. 2006; Rizzuto and Pozzan 2006; de Brito and Scorrano 2008; Hayashi et al. 2009). For example, within these microdomains, Ca\(^{2+}\) released by the ER through ryanodine (RyR) and inositol triphosphate (IP3) receptors moves down the steep potential gradient (~ −200 mV) across the inner mitochondrial membrane into the matrix, through a pore called the mitochondrial Ca\(^{2+}\) uniporter (MCU) (Kirichok et al. 2004). The MCU has recently been cloned, opening an important door for detailed study of its role and regulation (De Stefani et al. 2011).

Although MAMs create a means by which mitochondria can assist the ER in Ca\(^{2+}\) buffering, it seems highly likely that these junctions serve another signaling function that is independent of buffering per se. One well-documented

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function of Ca\textsuperscript{2+} flux through the MCU is stimulation of the tricarboxylic acid (TCA) cycle enzymes that produce reducing equivalents for oxidative phosphorylation (OXPHOS) (McCormack et al. 1990). In muscle, Ca\textsuperscript{2+} efflux from the ER stimulates contraction, which results in ATP hydrolysis, and in parallel stimulates mitochondrial OXPHOS, thereby helping to meet this metabolic demand (Jouaville et al. 1999). For this system to work effectively in neurons, the Ca\textsuperscript{2+} signal reaching mitochondria should accurately reflect metabolic needs associated with excitability. Recent work (Cardenas et al. 2010) shows that in neurons this system plays an essential role in regulating mitochondrial metabolism. However, because superoxide production is elevated by stimulation of the electron transport chain in the absence of sufficient ADP to drive the proton current through complex V (Korshunov et al. 1997), this system could have deleterious consequences if the “demand” on ATP does not meet the “supply” provided by Ca\textsuperscript{2+}-mediated acceleration of the TCA cycle. Interestingly, Foskett’s group has shown that elevated IP3R-mediated release of ER Ca\textsuperscript{2+} significantly increases the production of ROS by mitochondria (Muller et al. 2011). ER-stimulated production of mitochondrial ROS production could prove to be the driving force in PD pathogenesis (see below).

**NEURONAL PACEMAKING POSES A SPECIAL CHALLENGE TO IONIC HOMEOSTASIS**

In most neurons, the challenge to ionic homeostasis created by spiking is episodic. Striatal spiny projection neurons, for example, reside very near the K\textsuperscript{+} equilibrium potential some 40 mV away from spike threshold in the absence of synaptic potentials (Gerfen and Surmeier 2011). Excitatory synaptic input drives the membrane potential to spike threshold for hundreds of milliseconds at a time, but this activity is intermittent.

There is a subset of neurons in the brain that cannot stay quiet, spiking frequently and autonomously; that is, in the absence of synaptic input they spike on their own. Typically, autonomously generated spiking is very regular (synaptic input decreases this regularity); as a consequence, these neurons are often referred to as autonomous pacemakers. Autonomous pacemakers can be divided into fast and slow spiking classes. Fast autonomous pacemakers, such as globus pallidus or substantia nigra pars reticulata (SNr) GABAergic neurons, spike at rates above 20 Hz. At physiological temperature (Atherton and Bevan 2005; Chan et al. 2011) these neurons have very brief action potentials (<1 ms) and robustly express Ca\textsuperscript{2+}-binding proteins (Lee and Tepper 2007; Gross et al. 2011). Typically, autonomous spiking in these cells is driven by a combination of monovalent cation channels, such as Na\textsubscript{1.9} Na\textsuperscript{+} and hyperpolarization and nucleotide-activated cation (HCN) channels (Chan et al. 2004; Mercer et al. 2007). Slow pacemakers spike at much lower frequencies (~2–5 Hz) and have broad spikes (>2 ms). Neurons of this class are found scattered in many regions of the brain, including the hypothalamus, striatum, mesencephalon, and medulla (Paladini et al. 2007; Ding et al. 2010; Guzman et al. 2010). Although there has been much less work on pacemaking mechanisms in this class of neuron, it has been shown that their pacemaking typically depends on Na\textsubscript{1.9}, HCN channels, as well as cationic leak channels (Bean 2007). In a subset of these neurons, voltage-dependent L-type Ca\textsuperscript{2+} channels contribute to the inward current that drives pacemaking (Puopolo et al. 2007; Guzman et al. 2009; Putzier et al. 2009). These L-type channels have a distinctive pore-forming subunit (Ca\textsubscript{1.3}) that is coded for by the CACNA1D gene (Chan et al. 2007), which confers on the channel the ability to open at relatively negative membrane potentials (Surmeier et al. 2010). These channels are relatively rare and most L-type channels found in the brain have a Ca\textsubscript{1.2} pore-forming subunit (coded for by the CACNA1C gene) (Sinnegger-Brauns et al. 2009); this means that in most neurons, L-type channels open only during periods of strong depolarization, like during a spike. Although it has not been directly measured, sustained pacemaking must come with a significant metabolic price tag. In these neurons, the ionic gradients underlying excitability are under
constant assault. This should be particularly true in slow pacemakers with broad action potentials that use Ca\(^{2+}\) channels to help push the plasma membrane toward spike threshold. As noted above, Ca\(^{2+}\) must be pumped across the plasma membrane against a very steep potential gradient. The high-affinity, low-capacity PMCA appears to require one molecule of ATP for every Ca\(^{2+}\) ion moved across the plasma membrane. Moreover, in cells of this class that lack significant expression of fixed Ca\(^{2+}\)-buffering proteins (such as SNc DA neurons), Ca\(^{2+}\) entering neurons most likely is pumped not only by the proteins in the plasma membrane but also by those of intracellular organelles (such as the ER), adding to their metabolic burden.

NEURONS CARRY OTHER METABOLIC WEIGHTS

Although maintaining the ionic gradients underlying excitability undoubtedly creates a metabolic burden, this is not the only one neurons must carry. Synaptic transmission poses a burden. There are two aspects of this process that should be considered. Neurons communicate with one another largely (not exclusively) by release of transmitter molecules. These transmitters act at either ionotropic or metabotropic receptors. Ionotropic receptors change the membrane potential of postsynaptic neurons by opening a pore that allows ions to flow down their electrochemical gradients. With the termination of synaptic transmission, the pore closes. Although the ionic fluxes produced by a single synaptic event are much smaller than those produced by a spike, they still dissipate the ionic gradients established by pumps/exchangers and therefore consume ATP. Glutamatergic synaptic transmission, which engages receptors that are permeable to Na\(^{+}\) and Ca\(^{2+}\) ions, can create a significant metabolic burden on neurons, particularly in pathological conditions (Nicholls et al. 2007). The energetic demands posed by metabotropic receptors are more difficult to gauge, as they rarely directly couple to ion channel gating.

Synaptic transmission poses an energetic burden for the presynaptic neuron as well. Establishing the high transmitter concentration gradient across the vesicular membrane needed for effective synaptic function requires ATP (McMahon and Nicholls 1991). In the case of monoaminergic neurons, maintaining cytosolic transmitter concentration at low levels also is important to prevent its oxidation and the generation of reactive quinones (Sulzer and Zecca 2000; Greenamyre and Hastings 2004). The exocytosis process is itself ATP dependent. It is surprising that the combination of a reactive transmitter (dopamine) and the demands of exocytosis seem not to create a significant oxidative stress on DA terminal (Choi et al. 2011). Synapses also are typically at great distance from the nucleus and principal protein synthesis machinery, requiring ATP-dependent axonal transport of proteins, vesicles, and mitochondria. Synaptic protein lifetimes appear to be, in general, short, adding to the burden of maintaining them. The metabolic load accompanying a few dozen synapses seems not to pose a significant problem for most neurons. However, some neurons, such as SNc DA neurons, maintain as many as several hundred thousand synapses (Matsuda et al. 2009), potentially creating a sustained and significant metabolic bill.

NEURONS DEPEND ON MITOCHONDRIA

Because of their high basal energy demand, neurons depend on mitochondria to supply the ATP necessary for survival. Under normal conditions, glucose is metabolized by neurons to produce ATP. Glycolysis itself results in the generation of 2 mol of ATP per mole of glucose converted to pyruvate. However, when the pyruvate is transported into the mitochondria and further oxidized by the TCA cycle and the electron transport chain, the total yield of ATP increases to 36 mol per mole of glucose. The dependence of a cell on mitochondrial oxidative phosphorylation can be evaluated by creating genetic knockout models involving deletion of genes that are required for mitochondrial function. Mitochondria contain DNA that encodes a set of proteins that are required for electron transport chain function; these genes are transcribed and translated in the mitochondrial...
The mitochondrial transcription factor TFAM is required for that process, and deletion of TFAM abolishes mitochondrial transcription and undermines the stability of mitochondrial DNA. Accordingly, deletion of TFAM in a cell leads to loss of mitochondrial DNA and the development of a progressive decline in mitochondrial ATP production, even though the mitochondrial mass may be increased. The deletion of TFAM in specific cell lineages of the mouse reveals that some cells survive and function normally without functional mitochondria, whereas other cells fare poorly. For example, in epidermal progenitor cells the loss of TFAM has no apparent effect on proliferation or differentiation responses (Baris et al. 2011). In contrast, deletion of TFAM in the mouse heart leads to a progressive cardiomyopathy associated with the loss in ATP production and eventual lethality (Hansson et al. 2004). Likewise, deletion of TFAM leads to neuronal loss. For example, deletion of TFAM in SNc DA neurons leads to respiratory chain deficiency, cell loss, and impaired motor function (Ekstrand et al. 2007). This observation is consistent with a large literature demonstrating that mitochondrial toxins taken up by dopaminergic neurons lead to their demise (Przedborski and Vila 2003; Przedborski et al. 2004).

During intermediary metabolism, reducing equivalents in the form of nicotinamide adenine dinucleotide (NADH), nicotinamide adenine dinucleotide phosphate (NADPH), and FADH2 are obtained from the oxidation of metabolic substrates. The mitochondrial electron transport chain transfers these reducing equivalents ultimately to molecular O2. The free energy change associated with the transfer of electrons down the chain is used to translocate protons from the matrix to the intermembrane space, generating an electrochemical gradient across the inner membrane (typically −180 mV). Proton translocation occurs at complexes I, III, and IV and the electrons are ultimately transferred to O2 at the terminal cytochrome oxidase (complex IV). The transmembrane gradient is used by the FoF1-ATP synthase (complex V) to generate ATP from ADP and inorganic phosphate (Pi). Hence, the dynamic regulation of the mitochondrial transmembrane potential (ΔΨm) is determined by the balance between the rate of electron transport and proton translocation (tending to augment the potential) and the rate of ATP synthesis (tending to dissipate it). Some additional dissipation of the potential occurs through the activity of coupled membrane transport systems, uncoupling proteins, or nonspecific membrane leaks.

**MITOCHONDRIA AND OXIDANT STRESS**

For more than 40 years it has been known that mitochondria can potentially generate reactive oxygen species (ROS) (Jensen 1966b). The transfer of four electrons to O2 at complex IV yields H2O, but a small percentage of the electrons are captured by O2 at proximal sites in the chain. The transfer of a single electron to O2 yields superoxide anion, a free radical. Sites implicated in that process include complexes I, II, III, and possibly certain dehydrogenases of the TCA cycle (Murphy 2009). These radicals are released into the matrix compartment where they can potentially interact with lipids, proteins, DNA, or with each other.

The factors regulating the generation of superoxide from the electron transport chain are not fully understood. Some observations suggest that mitochondrial membrane hyperpolarization can augment ROS generation. In complex I, reducing equivalents obtained from NADH are transferred to ubiquinone-generating ubiquinol, a membrane-soluble electron carrier that delivers a pair of electrons to complex III. Complex II links the TCA cycle to the electron transport chain by oxidizing succinate to fumarate, delivering the electrons to complex III via ubiquinol. In isolated mitochondria given succinate to supply electrons into complex II in the absence of ADP, the membrane potential increases. Complex I is reversible and under hyperpolarized conditions a retrograde electron flux occurs from ubiquinol to NADH. This phenomenon is associated with a large increase in ROS production from complex I (Votyakova and Reynolds 2001). It can be inhibited by rotenone, a drug that inhibits reverse electron transport into the
distal end of complex I. It can also be inhibited by modest decreases in membrane potential, as the reverse flux is only favored when the membrane is highly polarized. Although this mechanism has been cited as an important source of ROS generation by isolated mitochondria, its significance under physiological conditions is questionable because the conditions that promote it (high membrane potential, succinate energy supply, lack of ADP) rarely occur in the intact cell.

Other mechanisms of ROS production in the mitochondrial matrix may relate to the rate of electron flux, with high rates of activity correlating with increased ROS generation. In that regard, the stimulation of mitochondrial activity by high ADP production in the cell would increase the transfer of electrons down the chain, potentially increasing the likelihood that some electrons will be captured by O₂ before they reach cytochrome oxidase (Lee et al. 2001; Rigoulet et al. 2011). The generation of superoxide in the matrix is also enhanced in cells subjected to hyperoxic conditions, where an increased availability of O₂ in the matrix compartment can enhance the probability that it will capture an electron from the electron transport chain. Finally, genetic defects in the expression or structure of various electron transport complex subunits can enhance the generation of superoxide. For example, in humans who are heterozygous for the B, C, or D subunits of succinate dehydrogenase (complex II), some cells can undergo somatic cell loss of heterozygosity, leading to the deletion of that subunit from the cell (Baysal et al. 2000). The A subunit of complex II contains the succinate dehydrogenase activity, whereas the B–D subunits are responsible for transferring the electrons to ubiquinone. If the B, C, or D subunit is disrupted while the A subunit is still expressed, the electrons derived from succinate oxidation cannot enter the electron transport chain and become stranded on the flavin group. This condition enhances the nonspecific generation of superoxide, which appears to enhance the tendency to form tumors (Guzy et al. 2008). The complete loss of complex II function in a cell leads to an inhibition of electron transport coupled to proton extrusion, and thus a loss of OXPHOS. Survival of the cell then depends entirely on ATP production by glycolysis, resulting in the release of lactic acid. Ironically, in that state the mitochondria shift to become consumers of ATP, which is taken up from the cytosol and used to maintain the mitochondrial membrane potential through reverse operation of complex V.

To protect against the potential detrimental effects of matrix oxidant stress, a complex set of “antioxidant” systems are expressed in the mitochondria that act to degrade ROS. Manganese superoxide dismutates (MnSOD) is a nuclear-encoded gene that is transcribed on cytosolic ribosomes and then imported into the mitochondrial matrix where it acts to dismute a pair of superoxide anions into hydrogen peroxide (H₂O₂). H₂O₂ is subsequently degraded by glutathione peroxidase and peroxiredoxins III and V, which are also expressed in the matrix. A distinct pool of reduced glutathione (GSH) resides in the mitochondria and is used by the reductase systems to clear H₂O₂ and lipid hydroperoxides. The resulting oxidized glutathione (GSSG) is subsequently re-reduced to GSH by glutathione reductase, which uses NADPH. The resulting NADP⁺ is then re-reduced to NADPH by dehydrogenases in the TCA cycle, such as isocitrate dehydrogenase-2. Ideally, under physiological conditions the rate of superoxide production is balanced by the rate of clearance, preventing the accumulation of superoxide/H₂O₂, thereby protecting the mitochondria from the damaging effects of oxidants.

MITOCHONDRIAL HYPOTHESIS OF AGING

In the 1920s, Pearl (1928) noted a correlation between the metabolic rate of an organism and its lifespan, such that animals with lower basal rates of metabolism lived longer than those with higher rates. This idea became known as the “rate of living theory of aging” (Pearl 1928) that linked metabolic rates to lifespan. In 1954, Commoner and coworkers (1954) reported detection of oxygen free radical formation in cells, using a paramagnetic resonance absorption technique. The idea that cells can generate...
oxygen radicals led Harman (1956) to hypothesize that aging and degenerative diseases may be attributable to the damaging effects of free radicals on cellular contents and extracellular connective tissue. According to that theory, interventions that lessen the generation or enhance the detoxification of reactive oxygen molecules in an organism should lessen the manifestations of aging and enhance longevity. This “oxygen radical theory of aging” evolved into the “mitochondrial theory of aging” after it was recognized that the mitochondrial electron transport chain is an important source of superoxide generation in the cell (Jensen 1966a).

Although mitochondria possess a sophisticated system for detoxifying reactive oxygen species, some of the ROS may survive long enough to react with lipids, proteins, or DNA. The probability of sustaining oxidant-mediated damage increases when the rate of ROS generation accelerates in response to stress, disease, or in mitochondria that become dysfunctional or damaged with aging. ROS interaction with mitochondrial complexes can lead to damage and/or dysfunction, and in some cases this damage can augment ROS production at that site or at other sites located earlier in the chain. One target of oxidative damage is mitochondrial DNA (mtDNA), which encodes a set of essential subunits of the electron transport chain as well as the mitochondrial transfer RNAs (tRNAs) required for translation. ROS-mediated damage to mtDNA is hypothesized to be an important mechanism underlying mutation in the mitochondrial genome (Beckman and Ames 1998). In that regard, ROS-mediated damage to mtDNA may lead to the expression of mutant forms of the electron transport complexes, whose altered function leads to an augmentation of their ROS production. This “vicious cycle” further injures the mtDNA and other mitochondrial constituents, thus amplifying the dysfunction and ROS generation (Wallace 2010). Indeed, multiple observations support the idea that mtDNA mutations contribute to the aging process (Aguilaniu et al. 2005). Furthermore, a large body of work has linked the level of mitochondrial oxidant stress to the aging phenotype and organismal lifespan, using genetic overexpression, knockdown or knock-out of antioxidant enzymes in widely diverse species (Jang and Remmen 2009).

Although the “mitochondrial theory of aging” is appealing in its simplicity and teleology, findings from other studies appear to challenge that model. First, mice that are heterozygous for the mitochondrial superoxide dismutase (MnSOD) show a 50% decrease in antioxidant capacity and evidence of increased mtDNA damage, yet they fail to show a significant decrease in lifespan compared with wild-type mice (Van Remmen et al. 2003). Conversely, increases in MnSOD activity induced in a transgenic line by overexpression of MnSOD were associated with decreased evidence of lipid peroxidation and decreased age-related decline in ATP production, yet no change in lifespan or age-related pathology (Jang et al. 2009). In other studies, Schriner et al. (2005) generated transgenic mice with mitochondria-targeted expression of catalase and observed an increase in median and maximal lifespan of ~20%, in association with a decrease in the frequency of mitochondrial deletions in skeletal muscle. However, when backcrossed into the C57BL6 background this phenotype was diminished whereas the mitochondrial catalase expression was retained. Thioredoxin-2 (Trx2) is a thiol reductase expressed in the mitochondria along with its partner, thioredoxin-2 reductase (Holmgren and Sengupta 2010). Genetic deletion of Trx2 (Trx2−/−) leads to embryonic lethality, indicating the importance of this system for maintaining mitochondrial protein thiol redox balance. Heterozygous animals (Trx2+/−) survive and show the expected evidence of increased mitochondrial oxidative stress and dysfunction (Perez et al. 2008) yet these mice do not show any evidence of shortened lifespan (Perez et al. 2009). Interestingly, overexpression of thioredoxin 1, which is expressed in the cytosol, enhances survival in the earlier stages of life, but does not extend maximum lifespan in the C57BL6 genetic background (Perez et al. 2011). In comparing multiple genetic models affecting oxidant stress, Richardson and colleagues (Perez et al. 2009) found that only Cu,Zn-SOD deletion had an effect on lifespan, leading them to question
whether the oxidative stress theory of aging is dead.

Using an elegant approach to disrupt mitochondrial DNA integrity, Trifunovic et al. (2004) and Kujoth et al. (2005) generated mouse lines with a knockin of a mutant version of the mitochondrial DNA polymerase POLG. The knockin carries a point mutation within the catalytic region responsible for proofreading mtDNA, leading to disruption of that function. Mice homozygous for this mutation accumulate mtDNA mutations over time and experience a significant shortening of lifespan compared with wild-type mice in the same background. These “mutator mice” also show an aging phenotype that is reminiscent of the pattern seen in aging humans. However, mitochondrial ROS generation and the level of lipid peroxidation and DNA oxidation was not increased in comparison to wild-type animals, raising questions about the theory that defects in mtDNA will lead to the generation of additional ROS by the electron transport chain. Studies of mtDNA in the heterozygous POLG mutant mice reveal significant increases in mtDNA point mutations compared with wild-type mice, yet those mice do not show a shortened lifespan or phenotypic evidence of advanced aging (Vermulst et al. 2007). However, the homozygous mice do have shortened lifespans; this group advanced the idea that there was a threshold for mtDNA deletions (arising from point mutations) that led to accelerated aging.

These observations highlight the controversy regarding the relationships among mitochondrial oxidant stress, the integrity of mtDNA, and the regulation of lifespan in diverse species. Although further work is needed to clarify the +apparent discrepancies in this field, these studies do provide some insight into the processes that may regulate the demise of DA neurons in SNc of patients with PD. First, it is important to note the strong association between aging and the development of PD, suggesting that SNc DA neurons may age more rapidly than other cell types in the brain. In view of the above controversy, it seems likely that the genes and processes regulating longevity of the organism may be distinct from those that regulate the aging phenotype in specific cell types. In that regard it is conceivable that some cells, by virtue of their phenotype and function, may be uniquely sensitive to genetic or environmental factors that can threaten their long-term survival. Those features may explain the relationship between PD and aging, yet have no mechanistic connection to the genetic factors regulating longevity.

PHYSIOLOGICAL FEATURES OF NEURONS AT HIGH RISK IN PD

Although PD is a disease of neurons, not all neurons are affected in PD (Braak et al. 2003). The vast majority of the neurons in the brain show no signs of pathology, even in the late stages of the disease. What then distinguishes neurons at risk in PD?

What Makes SNc DA Neurons Vulnerable in PD?

The neurons with the best-documented vulnerability are SNc DA neurons. The cardinal motor symptoms of PD, including bradykinesia, rigidity, and resting tremor, are the first to appear clinically and are clearly linked to the degeneration and death of SNc DA neurons (Hornykiewicz 1966). The efficacy of the clinical gold-standard treatment of L-dopa, a DA precursor, is testament to the centrality of DA neurons in the motor symptoms of PD. Why do these neurons die? A widely held theory implicates DA itself in their death, suggesting that oxidation of cytosolic DA (and its metabolites) leads to the production of cytotoxic free radicals (Greenamyre and Hastings 2004). However, there are reasons to doubt whether this type of cellular stress alone is responsible for the loss of DA neurons in PD. For example, there is considerable regional variability in the vulnerability of DA neurons in PD, with some areas being devoid of pathological markers (Matzuk and Saper 1985; Kish et al. 1988; Saper et al. 1991; Ito et al. 1992; Damier et al. 1999). Moreover, L-dopa administration (which relieves symptoms by elevating DA levels in PD patients) does not
accelerate disease progression (Fahn 2005), suggesting that DA itself is not a significant source of reactive oxidative stress, at least in the short term. Sulzer and colleagues have recently reported that Ca\(^{2+}\) entry through L-type channels stimulates DA metabolism in SNC DA neurons, pushing cytosolic DA concentrations into a toxic range with L-dopa loading (Mosharov et al. 2009). However, taken together the available data does not argue that DA itself is likely to be the principal culprit in the disease.

What is distinctive about SNC DA neurons is their physiological phenotype. Adult SNC DA neurons are slow, broad spike, autonomous pacemakers lacking much intrinsic Ca\(^{2+}\) buffering capacity (Foehring et al. 2009). This pacemaking activity is believed to be important in maintaining ambient DA levels in regions that are innervated by these neurons, particularly the striatum (Romo and Schultz 1990). Although most neurons rely exclusively on monovalent cation channels to drive pacemaking, SNC DA neurons also engage ion channels that allow extracellular Ca\(^{2+}\) to enter the cytoplasm (Ping and Shepard 1996; Bonci et al. 1998; Puopolo et al. 2007) leading to elevated intracellular Ca\(^{2+}\) concentrations (Wilson and Callaway 2000; Chan et al. 2007).

In adult SNC dopaminergic neurons, the currents that flow through these channels are of sufficient magnitude to sustain a membrane potential oscillation when voltage-dependent Na\(^{+}\) channels are blocked with tetrodotoxin (Fig. 3) (Guzman et al. 2009). This oscillation also depends on small conductance Ca\(^{2+}\)-activated K\(^{+}\) (SK) channels (Ping and Shepard 1996, 1999). The presence of this slow oscillatory potential and the sensitivity of pacemaking to dihydropyridines (DHP) led to the conjecture that L-type channels were necessary for pacemaking in SNC DA neurons (Nedergaard et al. 1993; Chan et al. 2007; Puopolo et al. 2007). However, the concentrations of DHP antagonist required to stop pacemaking was three orders of magnitude above the equilibrium binding constant for Ca\(_{\text{v}1.3}\) channels, raising questions about the specificity of drug action (Surmeier et al. 2010). What made these experiments difficult to interpret was that DHPs are not channel blockers (they do not plug the pore), but rather are negative allosteric regulators or antagonists (Sinnegger-Brauns et al. 2009). Furthermore, the allosteric interaction of DHPs with the channel is the voltage dependent. The equilibrium binding data was taken from membrane preparations where the transmembrane potential was nominally zero mV, maximizing affinity. At more negative membrane potentials, the affinity of the DHPs for L-type channels falls. However, the membrane potential of SNC DA neurons is always in a range near that needed to maximize binding affinity because of their pacemaking phenotype. We have modeled the interaction between DHP binding and pacemaking using the modulated receptor assumption advanced by Bean (1984); these simulations clearly suggest that submicromolar DHP concentrations should result in near complete antagonism (Surmeier et al. 2010; Ilijic et al. 2011). Using two-photon laser scanning microscopy in brain slices, this model was tested by monitoring dendritic Ca\(^{2+}\) fluxes in SNC DA neurons, which are attributable to L-type channels, during pacemaking (Guzman et al. 2009); these studies showed that submicromolar concentrations of the DHP isradipine eliminated any detectable fluctuation in distal dendritic Ca\(^{2+}\) concentration without altering pacemaking frequency, providing a strong confirmation of the hypothesis that Ca\(_{\text{v}1.3}\) channels are not necessary for pacemaking. Simulations of pacemaking have shown how this might happen (Guzman et al. 2009); in the model, as inward current through Ca\(_{\text{v}1.3}\) channels falls, outward current through SK K\(^{+}\) channels falls in parallel, leading to essentially no change in net transmembrane current and the maintenance of pacemaking. That said, in the absence of Ca\(_{\text{v}1.3}\) channels, pacemaking becomes less robust and stops with partial antagonism of HCN channels; adult SNC DA neurons with intact Ca\(_{\text{v}1.3}\) channels are resistant to this manipulation (Chan et al. 2007; Guzman et al. 2009).

Ventral tegmental area (VTA) DA neurons, which also are slow pacemakers, diverge from SNC DA neurons in two potentially important respects of relevance to this discussion. First, the Ca\(_{\text{v}1.3}\) Ca\(^{2+}\) channel density is dramatically...
Figure 3. Cav1.3 L-type Ca\textsuperscript{2+} channels contribute to the excitability of SNc DA neurons. (A) Pacemaking firing of SNc DA neurons recorded in current-clamp whole cell configuration in mouse midbrain slices. Pharmacological blockade of sodium channels with tetrodotoxin (TTX) leaves an ongoing membrane potential oscillation. (B) Another example of SNc DA neuron in pacemaking mode synchronized to dendritic Ca\textsuperscript{2+} oscillations imaged from a distal dendrite using Fluo4 Ca\textsuperscript{2+} indicator. Following application of TTX, the membrane potential oscillation is associated with an ongoing Ca\textsuperscript{2+} oscillation attenuated following antagonism of the DHP-sensitive L-type channels with isradipine. (C) Representative pacemaking SNc DA neuron displaying Ca\textsuperscript{2+} transients from a proximal dendrite (∼30 μm away from the soma, blue trace), and following bath application of isradipine to block L-type channels. (D) The same as part C, with the difference that Ca\textsuperscript{2+} imaging was performed on a distal dendrite (∼80 μm away from the soma, red trace). Antagonism of L-type channels attenuates the underlying Ca\textsuperscript{2+} oscillation in distal dendrites, while in a proximal dendrite; there is a residual Ca\textsuperscript{2+} transient that might be dependent on other voltage-gated Ca\textsuperscript{2+} channels. These results can suggest a potential gradient of L-type channel expression that increases as you move farther away from the soma. Parts C and D also show that isradipine did not affect pacemaking automaticity, suggesting that Ca\textsuperscript{2+} influx through L-type channels can be attenuated without affecting the pacemaking firing of SNc DA neurons. (Data used in this figure have been adapted from Chan et al. 2007 and Guzman et al. 2010, with permission from the authors.)
lower than in SNc DA neurons (Chan et al. 2007; Guzman et al. 2010). Second, the expression of the Ca\(^{2+}\)-buffering protein calbindin is much higher in VTA DA neurons than it is in SNc DA neurons (German et al. 1992a). The combination of these two factors results in the absence of detectable oscillations in dendritic Ca\(^{2+}\) concentration during pacemaking in VTA DA neurons. This should also lower the metabolic load on these neurons (Guzman et al. 2010). The human homologs of these VTA neurons have a significantly lower risk of degeneration in PD (Kish et al. 1988; German et al. 1992a; Pignatelli et al. 2005; Belzung and Brunello 2003), linking metabolism to risk in PD at a different point in the metabolic spectrum. There are many unanswered questions, though, about how these physiological features are linked to risk. How, for example, do Ca\(^{2+}\)-buffering proteins shape the metabolic burden of Ca\(^{2+}\) influx? Based on experiments examining intrinsic Ca\(^{2+}\)-buffering capacity using whole cell patch pipettes, it appears that most intrinsic Ca\(^{2+}\)-buffering proteins are immobile, as they are not washed out by dialysis (Augustine et al. 2003). By binding Ca\(^{2+}\), these proteins “seques- ter” it away from signaling proteins and organelles during opening of membrane channels. When these channels close and cytosolic Ca\(^{2+}\) levels begin to decrease with activity of the pumps/exchangers, Ca\(^{2+}\) could move from this bound pool to become available for transport out of the cell. Because the interaction between Ca\(^{2+}\) and the binding proteins is ionic and does not involve ATP-dependent sequestration into an organelle, this could be an energetically “cheap” way for the cell to manage Ca\(^{2+}\) influx.

Recent work by our group has shown that maintained opening of L-type Ca\(^{2+}\) channels in SNC DA neurons creates a basal mitochondrial oxidant stress (Guzman et al. 2010). Recall that these neurons lack abundant Ca\(^{2+}\)-buffering proteins (Foehring et al. 2009). These studies used a transgenic mouse that expressed a mitochondrially targeted redox-sensitive variant of green-fluorescent protein (mito-roGFP) (Hanson et al. 2004; Guzman et al. 2010) expressed under control of the tyrosine hydroxylase (TH) promoter (Fig. 3). The use of roGFP allowed mitochondrial matrix redox state to be quantitatively estimated, something not possible with conventional redox probes. Using two-photon laser scanning microscopy to monitor mito-roGFP in brain slices from young adult mice, we found that the engagement of plasma membrane Ca\(_{\text{A,1.3}}\) L-type Ca\(^{2+}\) channels during normal autonomous pacemaking created an oxidant stress in the mitochondria that was specific to the vulnerable SNc DA neurons and not apparent in neighboring VTA DA neurons. The oxidant stress engaged defenses that induced transient, mild mitochondrial depolarization or uncoupling. The mild uncoupling was not affected by deletion of cyclophilin D, which is a component of the permeability transition pore, but was attenuated by genipin and purine nucleotides, which are antagonists of cloned uncoupling proteins. Knocking out DJ-1 (also known as PARK7 in humans and Park7 in mice), which is a gene associated with an early-onset form of PD, down-regulated the expression of two uncoupling proteins (UCP4 [SLC25A27] and UCP5 [SLC25A14]), compromised Ca\(^{2+}\)-induced uncoupling, and increased oxidation of matrix proteins specifically in SNc dopaminergic neurons. The results with the DJ-1 knockout—showing that the impact of DJ-1 deletion depends on a physiological phenotype that engages mitochondrial oxidant defenses—provides an example of how mutations in a widely expressed gene can affect a select subpopulation of neurons. That said, the mechanisms governing mitochondrial matrix redox state are complex and the hypothesis that UCP4/5 contribute to this regulation must be rigorously tested; the most obvious strategy would be to knock the mRNA coding for these proteins down using viral delivery of small interfering RNAs (siRNAs).

The metabolic demands posed by maintaining transmembrane ionic gradients underlying excitability led Nicholls (2002) to postulate that neurons are at risk in neurodegenerative disease because they have a modest bioenergetic or respiratory reserve. This reserve is defined as the difference between the maximum capacity for ATP generation by oxidative phosphorylation and the basal consumption of ATP. The smaller
this respiratory reserve, the more likely episodic demands on metabolism, such as exposure to a toxin or bursts of spiking, the more likely cellular ATP levels will decrease and create a bioenergetics crisis, leading to loss of membrane potential or other ATP-dependent processes. Persistent loss of membrane potential will lead to massive Ca\(^{2+}\) influx and cell death (Beal 1995; Choi et al. 2011). Clearly, slow pacemaking neurons with broad action potentials, sustained Ca\(^{2+}\) influx, and low intrinsic Ca\(^{2+}\)-buffering capacity, such as SNc DA neurons, would be at the bad end of the respiratory reserve distribution of neurons, putting them at risk for insults (Fig. 4A).

But if this type of neuronal design puts neurons at risk, why wouldn’t evolution have eliminated it? It appears that most of the neuronal systems that have adopted this phenotype are very tolerant of cell loss. For example, roughly three-quarters of the SNc DA neuron population must be lost before the motor symptoms are severe enough to prompt a visit to the neurologist (Zigmond et al. 1990). This takes, on average, six decades to occur, which is well past the reproductive period and well past the normal life expectancy until recently. As a consequence, there could not be any evolutionary pressure to change this design.

Is this design merely an accident or unintended consequence of a larger genetic program controlling the SNc DA neuron phenotype? There are some reasons to believe that it is not an accident but an adaptation with unintended consequences. Although not necessary, L-type Ca\(^{2+}\) channel currents do increase the robustness of pacemaking in SNc DA neurons (Guzman et al. 2009); as pacemaking is critical to maintaining normal basal ganglia function, designing a redundant, fail-safe system makes some sense. Dendritic L-type Ca\(^{2+}\) channels also have been shown to boost the frequency

**Figure 4.** Neurons with the pacemaking phenotype might have a diminished respiratory reserve, putting them at risk of bioenergetics failure. (A) Schematic model of the respiratory capacity of a quiescent neuron, an active neuron, and an active Ca\(^{2+}\)-regulated neuron. Respiratory capacity here is defined as the sum of the basal respiration of sustained cellular function and the respiratory reserve, which represents the “back-up” extra fuel for the cell that is normally used under active conditions (e.g., an excitable neuron). Active neurons with a Ca\(^{2+}\)-handling phenotype like SNc DA neurons, reach a bioenergetics cliff owing to the enhanced stimulation of the tricarboxylic Krebs cycle (TCA) by Ca\(^{2+}\). This boost increases respiratory capacity of the cell to generate and supply more ATP necessary to support further Ca\(^{2+}\) extrusion. (B) Hypothetical plot of ROS production by the electron transport chain (ETC) as a function of inner mitochondrial membrane potential. Also plotted are the effects of Ca\(^{2+}\) stimulation of the TCA cycle and the effects of complex V and uncoupling proteins (UCPs). (C) Metabolic demand expressed as an increase in respiratory capacity comes at a trade-off of generating reactive oxygen species (ROS) (shown in the graph as the increase in oxidant stress), which will feedback to negatively affect proteostatic function and compromise cellular respiration until it reaches the point of a bioenergetics failure, in which a cell can no longer meet the metabolic demand and will fail to continuously supply the cell with energy fuel to extrude Ca\(^{2+}\).
of synaptically driven spiking in SNc DA neurons (Wilson and Callaway 2000; Medvedev et al. 2003); although this enhancement in “burst” spiking is modest, it could elevate DA release just enough to have conferred some survival value by facilitating striatal motor learning. Another possibility is that the expression of Ca\(^{2+}\) channels is an outgrowth of the modest respiratory reserve conferred by the pacemaking phenotype (Fig. 4B). Ca\(^{2+}\) entry into the mitochondrial matrix accelerates the TCA cycle and increases respiratory capacity. Denton and McCormack (1986) showed that the activities of pyruvate dehydrogenase, isocitrate dehydrogenase-3 (IDH3) and 2-oxoglutarate dehydrogenase are enhanced by low micromolar concentrations of Ca\(^{2+}\), so the uptake of cytosolic Ca\(^{2+}\) into the matrix is likely to enhance the generation of reducing equivalents that feed into the electron transport chain. If this increase in capacity exceeds the cost associated with Ca\(^{2+}\) extrusion, then it would increase the respiratory reserve and the resilience to insults and transient stress. This is a strategy used by muscle to deal with the metabolic burden created by contraction. The trick is that supply and demand needs to be matched. Accelerating the TCA and the supply of electrons to the ETC in the absence of increased ATP demand leads to hyperpolarization of the inner mitochondrial membrane (IMM) and increased ROS production (Brand et al. 2004a,b). Matching is straightforward in muscle because the ER orchestrates both the increased supply (increased respiration) and the increased demand (contraction). But this is not the case in neurons (at least it is not obvious). If anything, ER release of Ca\(^{2+}\) is associated with activation of Ca\(^{2+}\)-dependent K\(^{+}\) channels and decreased spiking (Morikawa et al. 2003). Although plausible and consistent with much of what is known about metabolic regulation in neurons, our understanding of these processes is cursory at best. Much more needs to be done in intact neurons that generate normal patterns of activity to determine whether this hypothesis has merit.

Can oxidant stress (regardless of how it is generated) and mtDNA damage induced by oxidants explain the selective loss of SNc DA neurons in PD? Deletion mutations in mtDNA can arise when H\(_2\)O\(_2\) in the matrix introduces double-strand breaks; accordingly the frequency of these mutations is decreased in hearts of mice expressing mitochondrial catalase (Vermulst et al. 2008). In two studies published simultaneously, Bender et al. (2006) and Kraytsberg et al. (2006) assessed the abundance of mitochondrial DNA deletions, as opposed to point mutations, in SNc neurons from human subjects. Compared to younger subjects, the number of mtDNA deletions was significantly greater in SNc neurons from older subjects. In contrast, undetectable levels of deletions were found in cerebral cortex, cerebellum, or dentate nucleus of aged individuals (Kraytsberg et al. 2006). This indicates that cell-specific differences in the occurrence of deletions can exist, consistent with the cell-specific manifestations of PD. Comparisons among single neurons from the same subject revealed that one neuron might contain no deletions, whereas another would contain multiple copies of a single species—indicating that they originated from a single initial mutant DNA copy that was clonally amplified in that cell (Kraytsberg et al. 2006). The mechanisms responsible for the clonal expansion of a single deletion mutant copy of mtDNA are not known. But if ongoing accumulation of random mutational events had been occurring, one would expect to find a heterogeneous mixture of point mutations in a given cell. Instead, it appears that distinct somatic deletion mutations have developed with age, occurring independently in multiple cells of SN, leading to the appearance of a heterogeneous patchwork of cells each carrying its own pattern of mtDNA deletion.

The point mutations that accumulate in the polymerase-γ (POLG) mutant mice can shorten lifespan when they exceed a critical level (Kujoth et al. 2005), but there is no evidence to suggest that either the homozygous or the heterozygous mice develop motor control defects as they age, or that they develop excessive oxidant stress as a consequence of these mutations. In contrast, normal SNc neurons do show enhanced mitochondrial oxidant stress by virtue of their Ca\(^{2+}\) handling, and the formation of...
H$_2$O$_2$ can induce deletion mutations in mtDNA. These observations suggest that the accumulation of point mutations leading to an augmentation of mitochondrial oxidant stress is unlikely to explain the demise of SNc neurons in PD.

An unresolved question is whether the induction of deletion mutations in mtDNA of SNc neurons can lead to an amplification of the ROS generation and trigger the "vicious cycle" hypothesis within this selective population of cells. Deletions of mtDNA can lead to the expression of truncation mutant proteins, or to the complete loss of subunit expression. When one subunit of a mitochondrial complex is genetically deleted, the remaining subunits are still expressed and degraded, which can lead to the generation of an unfolded protein response in the matrix (Zara et al. 2007). For example, Andrew Dillin and colleagues (Durieux et al. 2011) find that deletion of cytochrome oxidase in worm neurons leads to a mitochondrial stress response in the organelle that is transmitted to other tissues and affects lifespan. By augmenting mitochondrial, and possibly ER stress responses, deletion mutations in mtDNA can potentially amplify oxidant stress in mitochondria and other cellular compartments, pushing the cell that normally deals with enhanced oxidant stress even closer to the edge. Hence, the cellular consequences of mtDNA damage may depend importantly on the nature of the mutation.

Deletion mutations in mtDNA can potentially augment mitochondrial ROS production by simply blocking the electron transport chain, which then halts the TCA cycle. Complete loss of any protein encoded by a mitochondrial gene is sufficient to abolish electron transport chain function. In that situation, flavin groups, iron-sulfur clusters, and heme proteins in the more proximal complexes of the chain remain fully reduced. However, electrons can still escape through the generation of superoxide, augmenting oxidant stress. At the same time, mitochondrial antioxidant capacity is undermined because the loss of electron transport causes the TCA cycle to halt, preventing the generation of NADPH. The scavenging of H$_2$O$_2$ by glutathione peroxidases and peroxiredoxins depends on the presence of reduced glutathione (GSH), which is required by the reductases that sustain their activity. Likewise, the reduction of oxidized mitochondrial protein thiols by glutaredoxin and Trx2 requires GSH. The oxidized glutathione generated by these reductase systems is re-reduced to GSH using NADPH generated by the TCA cycle. Hence, the loss of a single mitochondrial gene can augment mitochondrial ROS generation and simultaneously undermine the ability of mitochondria to deal with that stress by limiting the supply of NADPH needed for antioxidant functions.

Mitochondrial DNA deletion mutations can also affect cell survival through a bioenergetic mechanism. When the abundance of damaged mtDNA is below a critical level, the normal copies of mtDNA can supply the organelle with adequate levels of the encoded proteins to maintain function and ATP production. However, when the abundance of the mutant form exceeds a critical level (typically 60%), the phenotypic defect in function becomes evident. Accordingly, Kraytsberg et al. (2006) found that the presence of mtDNA deletions correlated strongly with the absence of cytochrome oxidase immunostaining (Bender et al. 2006). Mitochondrial DNA encodes three critical subunits of cytochrome oxidase, so these findings suggest that the accumulation of mtDNA deletions is responsible for the development of respiratory insufficiency in the affected cells. DA neurons of SNc depend on mitochondrial DNA for survival (Ekstrand et al. 2007), so the accumulation of these mutations can potentially lead to the development of a bioenergetic deficiency that becomes lethal over time. As discussed previously, loss of functionality in the electron transport chain shifts the mitochondria to become ATP consumers, further stressing the bioenergetic status of the cell. In PD, it is conceivable that the progressive clonal expansion of the deletion mutant could occur over many years, resulting in the progressive demise of DA cells.

How do SNc neurons in aged subjects compare with those from individuals with PD? Bender et al. (2006) found that the degree of
mtDNA deletions was somewhat higher in neurons from affected individuals compared with aged control subjects. This was associated with a significantly greater proportion of cytochrome oxidase-deficient cells, compatible with the idea that bioenergetic crisis could be responsible for the progressive cellular loss. Consistent with the study of Kraytsberg et al. (2006), this group detected clonal expansion of the unique species of mtDNA deletions in individual SNc neurons, indicating that these are indeed somatic mutations. In contrast, high levels of deleted mtDNA were not detected in hippocampus.

**What about Non-DA Neurons in PD?**

Many lines of study ranging from histological analysis of PD brains to clinical examination of PD patients have made it clear that pathogenesis is not limited to SNc DA neurons. Based on LB/LN pathology, in the early and middle stages of the disease, many small groups of neurons in the brain stem, diencephalon, and telencephalon are affected. Foremost among these are neurons in the dorsal motor nucleus of the vagus (DMV), in the locus coeruleus, in the raphe nuclei, in the gigantocellularis nucleus, in the tuberomamillary nucleus of the hypothalamus, in the olfactory bulb, and in the basal forebrain (German et al. 1992b; Del Tredici and Braak 2004).

Most of these neurons have not been studied in detail, but enough is known to draw a couple of conclusions. First, all of these nuclei are dominated by neurons that are spontaneously active (Williams et al. 1984; Chan and Chan 1989; McCann and Rogers 1990; Serafin et al. 1990; Travagli et al. 1991; Aston-Jones 2005). Second, many of these neurons have broad action potentials. Much less is known about whether these neurons share the other phenotypic features of SNc DA neurons that are likely to determine vulnerability, such as prominent Ca²⁺ currents. What is known is consistent with this hypothesis. For example, raphe neurons show pronounced Ca²⁺-dependent pacemaking potentials (Burkli and Aghajanian 1987). Both locus coeruleus and tuberomamillary neurons are autonomous pacemakers that engage L-type Ca²⁺ channels (Williams et al. 1984; Stevens and Haas 1996; Taddese and Bean 2002). Our unpublished work also has found that locus coeruleus neurons also have low intrinsic Ca²⁺-buffering capacity and basal mitochondrial oxidant stress that is sensitive to L-type channel antagonism. DA neurons in the olfactory bulb also are autonomous pacemakers and rely on Ca²⁺ channels (although not L-type channels) (Pignatelli et al. 2005; Puopolo et al. 2005). Although olfactory deficits have been associated with PD (Postuma et al. 2006), there is no obvious loss of olfactory bulb DA neurons (Huisman et al. 2008). Although this would seem to run counter to the phenotype hypothesis, this could simply be a consequence of the capacity of this region for adult neurogenesis (Pignatelli et al. 2009). Thus, a reasonable hypothesis is that all of the neurons displaying a strong risk in PD share a common physiological phenotype.

**RECONCILIATION WITH CURRENT MODELS OF PD PATHOGENESIS**

The proposition that a neuronal phenotype leads to excessive mitochondrial oxidant stress, mitochondrial DNA deletion, and bioenergetic deficiency is consistent with a large body of evidence implicating these organelles in PD pathogenesis (Schapira 2011; Trancikova et al. 2011). The following are important questions in that model that still need to be addressed: Does the oxidant stress extend from the mitochondria to other parts of the cell? Is oxidant stress responsible for the demise of the dopaminergic SNc cells in humans? What are the targets of the oxidant stress that lead to that demise, and what mechanisms link the neuronal phenotype to the long latent period before the onset of disease symptoms?

Another prominent theory in the field is that PD is essentially a protein misfolding or proteostatic disorder (Bosco et al. 2011). The presence of Lewy bodies and Lewy neurites in substantia nigra is a common finding in patients with PD; the major filamentous protein component of the LBs and LNs is α-synuclein (Spillantini et al. 1998). Mutations in the α-synuclein
inherited PD associated with the accumulation of α-synuclein aggregates in humans, providing a strong link to the disease (Polymeropoulos et al. 1997; Kruger et al. 1998; Dawson and Dawson 2003a,b; Berg et al. 2005). Accumulation of misfolded proteins can occur when an imbalance develops between the factors tending to promote protein unfolding (oxidant stress, pH stress, thermal shock, prion-mediated disruption) and the systems responsible for removal (ubiquitylation, proteasomes) or refolding (molecular chaperones). It is therefore reasonable to expect that mutations promoting misfolding of an abundantly expressed protein can lead to a disruption of this balance and the accumulation of aggregates, with functional consequences for cell survival. Indeed the expression of mutant versions of α-synuclein in flies leads to the development of motor control defects and inclusion bodies that resemble PD in humans (Fean and Bender 2000). However, in patients with PD who lack such mutations, it is not clear whether the cell death is caused by the accumulation of these aggregates.

Deficits in proteasome clearance of misfolded proteins and autophagy have unequivocally been linked to neurodegeneration in animal models of PD (Shastry 2003). In some cases the cellular degeneration may result from a nonspecific accumulation of disordered proteins such as α-synuclein, but in other cases it can develop because a specific functional protein is not cleared normally. In that regard, genetic defects in Parkin, an E3 ubiquitin ligase, have been linked to human PD (Shimura et al. 2000). A protein target of Parkin is PARIS, which functions as a transcriptional repressor of peroxisome proliferator-activated receptor γ (PPARγ) cofactor-1α (PGC-1α) (Shin et al. 2011). PGC-1α is a major regulator of mitochondrial biogenesis (Lin et al. 2002), so suppression of its activity by PARIS, which accumulates in the absence of Parkin, could drive cells toward bioenergetic deficiency by limiting the number of mitochondria. Indeed, deletion of Parkin from SN led to progressive loss of dopaminergic neurons in a PARIS-dependent manner (Shin et al. 2011).

In the case of α-synuclein, it is not known whether the aggregation of wild-type α-synuclein in the aging human is a proximal initiator of PD, whether it contributes to the cell death as a distal effector, or whether it represents a marker of a cell already committed to a terminal pathway. Some evidence suggests that oxidation of methionine residues in α-synuclein (which lacks cysteine residues) may promote its accumulation as a stable oligomer while preventing the fibrillation of unmodified α-synuclein (for review see Glaser et al. 2005). In view of our data linking the cellular Ca\(^{2+}\) phenotype to the generation of oxidant stress, it is conceivable that the oxidative modification of α-synuclein could lead to its subsequent oligomerization in dopaminergic SNc cells. A number of studies have provided evidence that elevated Ca\(^{2+}\) concentration promotes aggregation of α-synuclein (Gomez-Tortosa et al. 2001; Nielsen et al. 2001; Lowe et al. 2004; Tamamizu-Kato et al. 2006; Esteves et al. 2010; Fortin et al. 2010; Nath et al. 2011). It is also possible that α-synuclein toxicity compromises Ca\(^{2+}\) homeostasis, particularly in neurons like SNc DA neurons where Ca\(^{2+}\) poses a challenge normally (Mattson 2007; Hettiarachchi et al. 2009; Belal et al. 2011; Vekrellis et al. 2011). As the oligomers of α-synuclein may be cytotoxic, this mechanism furthers the notion that the phenotype of these neurons, their Ca\(^{2+}\)-dependent ROS generation, and their tendency to express a protein that is vulnerable to forming toxic aggregates when oxidized, may explain the cell-specific manifestations of the disease. Furthermore, the clearance of misfolded proteins requires energy in the form of ATP, so the bioenergetic consequences of mitochondrial dysfunction (discussed earlier) could further compromise the clearance of misfolded proteins, leading to their accumulation in protein dumps that grow to become LBs and LNs (Esteves et al. 2011).

ARE L-TYPE Ca\(^{2+}\) CHANNELS A VIABLE THERAPEUTIC TARGET?

Although there are several factors governing the loss of SNc DA neurons, most of them cannot be manipulated. The exception is the engagement of
L-type Ca^{2+} channels. These channels are antagonized by orally deliverable dihydropyridines with good brain bioavailability that have a long record of safe use in humans (Yamada et al. 1990; German et al. 1992a; Chan et al. 2007; Guzman et al. 2010; Ilijic et al. 2011). SNc DA neurons that express the Ca^{2+}-binding protein calbindin have a diminished sensitivity to toxins and to PD (Yamada et al. 1990; German et al. 1992a).

Is there evidence that DHP use might work in humans to prevent or slow PD? Ca^{2+} channel antagonists (CCAs), including the DHPs used in animal studies, are commonly used in clinical practice to treat hypertension, creating a potential database to be mined. A case-control study of hypertensive patients found a significant reduction in the observed risk of PD with CCA use, but not with medications that reduce blood pressure in other ways (Becker et al. 2008). More recently, a large Danish data set has been examined (Ritz et al. 2010). The investigators agreed with the main conclusions of the Becker et al. study but extended their findings by showing that only DHPs that cross the blood–brain barrier (BBB) are associated with reduced PD risk (~30%). Given the short period of treatment in many cases (~2 yr), variable dosing, and low relative affinity of DHPs for Ca_{1.3} Ca^{2+} channels (compared to Ca_{1.2} channels (Mannhold et al. 1995; Kupsch et al. 1996; Eisenberg et al. 2004), this is a surprisingly strong association and lends further credence to the proposition that a BBB-permeable and potent Ca_{1.3} antagonist could be a very effective neuroprotective agent. It should be noted that a small, recent study (Simon et al. 2010) failed to find a significant relationship between use of Ca^{2+} channel antagonists and PD risk; however, this study did not take into account the type of Ca^{2+} channel antagonist used or the pharmacokinetic properties of the compounds. This undermines any conclusion to be drawn from the study. After all, Ritz et al. (2010) showed that even amlodipine—a DHP with reasonable affinity for Ca_{1} channels—does not reduce the observed risk of developing PD, but amlodipine does not cross the blood–brain barrier.

Although strongly suggestive, epidemiological studies are not a substitute for a prospective controlled clinical trial. In the absence of a selective Ca_{1.3} Ca^{2+} channel antagonist, the DHP isradipine is the most attractive drug for such a trial. Isradipine has a relatively higher affinity for Ca_{1.3} Ca^{2+} channels than the other known DHP and has good brain bioavailability (Koschak et al. 2001; Scholze et al. 2001). At the doses used to treat hypertension, isradipine has relatively minor side effects (Fitzon and Benfield 1990). The question is whether it will prove neuroprotective at doses tolerated by the general population. Pharmacokinetic studies by our group have found that plasma concentrations of isradipine achieved in mice that are protected against systemic 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) administration are very close to those achieved in humans with a very well tolerated daily dose of isradipine (~2–4 ng/mL in mice; 1–2 ng/mL in humans at 10 mg/d, Dynacirc CR), suggesting that neuroprotection is achievable. A more recent study using an intrastriatal 6-hydroxydopamine model has shown that systemic administration of isradipine produces a dose-dependent protection of both SNc DA axon terminals and cell bodies at plasma concentrations in a similar range (IC50 ~4–8 ng/mL) (Ilijic et al. 2011). The plasma concentrations required for significant protection against this acute challenge were greater than those needed in the chronic MPTP model, but were are still near the range achievable in humans. It is important to note that these studies suggest that protection is afforded by partial antagonism of Ca_{1.3} channels, minimizing any complications that might attend near complete disruption of these channels. Using a modulated receptor model derived from an early paper by Bean (1984), Ilijic et al. estimated that ~40%–60% of the Ca_{1.3} channels were antagonized at the half-maximal dose for protection from intrastriatal 6-OHDA injection. Given the slow progression of PD, it is highly likely that a more modest antagonism would suffice in humans.

The ideal candidate for DHP therapy would be in the very early stages of SNc loss, before the onset of symptoms. Unfortunately, there are no biomarkers that would allow the identification of presymptomatic PD patients. As a
consequence, the most likely subjects for a clinical trial are those that have been recently diagnosed with PD. In these early-stage patients, SNc DA cell loss is substantial (>60%) and the remaining neurons might be compromised in ways not seen in healthy tissue. Nevertheless, disease progression could be tracked in a longitudinal study to determine whether treated patients show a slower rate of motor deficit progression. There is growing evidence that inflammation could have an important impact on disease progression at this stage (Hirsch and Hunot 2009). Antagonism of L-type channels might be helpful but not enough to significantly slow progression if this is the case.

It is also worth considering how DHPs compare with other drugs that are being tested in clinical neuroprotection trials for PD. Although early trials with creatine, coenzyme Q10, and the mitochondrial antioxidant compound MitoQ have been disappointing (Shults et al. 1997), this is not sufficient evidence to reject the idea that oxidative stress contributes to PD. Coenzyme Q10 is an electron acceptor for complexes I and II that appears compromised in PD patients (Shults et al. 1997) and is neuroprotective in animal models of PD (Beal et al. 1998). MitoQ is a mitochondria-targeted version of coenzyme Q10 that functions as a scavenger of mitochondrial oxidants (Smith and Murphy 2011). Creatine is a substrate for ATP production that can both improve mitochondrial efficiency and reduce oxidative stress by buffering fluctuations in cellular energy production (Matthews et al. 1999). One possibility is that antioxidant treatment initiated after the point of clinical diagnosis is ineffective because the cells have already accumulated a critical level of damage. By analogy, myocardial damage in ischemia-reperfusion is mediated by excessive oxidant stress, yet antioxidants given in the days after myocardial infarction are ineffective at rescuing cell function because the oxidant damage is already done. It is also important to consider that the effect of antioxidants will depend on where the oxidant stress is generated and where the critical targets of these oxidants reside. Clearly, an antioxidant distributed, for example, in membranes could be minimally effective in scavenging oxidants in the mitochondrial matrix. If antioxidant therapy by itself is ineffective, then approaches aimed at improving mitochondrial function rather than attacking the source of stress on mitochondria could be more effective. For example, deprenyl could prove to have neuroprotective effects by virtue not of its ability to inhibit the degradation of DA, but by its ability to induce the expression of antioxidant defenses (Magyar and Szende 2004). Because their sites of action differ within the chain of events leading to oxidative stress and mitochondrial dysfunction, a combination therapy could prove more effective than any one alone.

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# Physiological Phenotype and Vulnerability in Parkinson's Disease

D. James Surmeier, Jaime N. Guzman, Javier Sanchez and Paul T. Schumacker

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