Towards an Understanding of Energy Impairment in Huntington’s Disease Brain

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Abstract. This review systematically examines the evidence for shifts in flux through energy generating biochemical pathways in Huntington’s disease (HD) brains from humans and model systems. Compromise of the electron transport chain (ETC) appears not to be the primary or earliest metabolic change in HD pathogenesis. Rather, compromise of glucose uptake facilitates glucose flux through glycolysis and may possibly decrease flux through the pentose phosphate pathway (PPP), limiting subsequent NADPH and GSH production needed for antioxidant protection. As a result, oxidative damage to key glycolytic and tricarboxylic acid (TCA) cycle enzymes further restricts energy production so that while basal needs may be met through oxidative phosphorylation, those of excessive stimulation cannot. Energy production may also be compromised by deficits in mitochondrial biogenesis, dynamics or trafficking. Restrictions on energy production may be compensated for by glutamate oxidation and/or stimulation of fatty acid oxidation. Transcriptional dysregulation generated by mutant huntingtin also contributes to energetic disruption at specific enzymatic steps. Many of the alterations in metabolic substrates and enzymes may derive from normal regulatory feedback mechanisms and appear oscillatory. Fine temporal sequencing of the shifts in metabolic flux and transcriptional and expression changes associated with mutant huntingtin expression remain largely unexplored and may be model dependent. Differences in disease progression among HD model systems at the time of experimentation and their varying states of metabolic compensation may explain conflicting reports in the literature. Progressive shifts in metabolic flux represent homeostatic compensatory mechanisms that maintain the model organism through presymptomatic and symptomatic stages.

Keywords: Huntington’s Disease, homeostasis, intermediate metabolism, flux analysis, mitochondrial dysfunction

ABBREVIATIONS

Aco aconitase
AAT aspartate amino transferase
αKG α-ketoglutarate
ALAT alanine amino transferase
AMPK AMP kinase
ANT adenosine nucleotide translocator
Aralar glutamate-aspartate transporter
Asp aspartate
Carn carnitine
CAT I carnitine acyltransferase I
CAT II carnitine acyltransferase II
CI complex I
CII complex II, SDH
CIII complex III
CIV complex IV, cytochrome oxidase
CK creatine kinase
CMRO₂ cerebral metabolic rate of oxygen consumption
COX cytochrome oxidase
Cr creatine
CS citrate synthase

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INTRODUCTION

Energy impairment is an acknowledged part of the Huntington’s disease (HD) phenotype. Yet within the CNS, how and why metabolic changes occur remain controversial. Mechanistically, abnormal energy metabolism in HD results from multiple mechanisms including altered mitochondrial axonal transport, dynamics and biogenesis due to nuclear transcriptional disruption [1–3]. Beyond issues of the size and distribution of mitochondrial mass, controversies remain concerning the existence of abnormalities or imbalances in the flux of metabolites through the various energy generating pathways to produce adequate ATP. Additionally, altered flux may reflect abnormal protein regulation or energy demands. Much attention has been focused upon mitochondrial function, specifically the electron transport chain (ETC), while deficits in glycolysis, the tricarboxylic acid cycle (TCA) and fatty acid oxidation (FAO) have received less attention.

Three different frameworks can be proposed to explain the metabolic dysfunction in HD brains. The first posits that mitochondrial biogenesis, dynamics, trafficking and mitophagy are altered resulting in a dearth of energy generating organelles in appropriate synaptic sites. Each of these mechanisms has been demonstrated in a variety of HD model systems [1–5]. However the presence of these restrictions on mitochondria does not explain how the organism grows initially and continues to survive. The second proposes that the energy generating enzyme complexes in the ETC of CNS mitochondria are compromised due to deficits in expression or functionality. Evidence for alterations in oxidative metabolism comes from post mortem human tissue and mouse models towards the end of their lives [6–8].
metabolic enzymes. Transcriptional dysregulation oxidative damage may progressively inactivate some phorylation on a temporal basis. Slowly accumulating mass action drives shifts in the balance changes may follow a similar contextual and temporal components [12]. Metabolic mitochondrial functions. Electrophysiological changes in effects should influence how multi stresses mitochondrial demise that ensues [11]. Similar contextual model systems influences the mechanisms of neuronal demise that ensues [11]. Similar contextual effects should influence how muh1 stresses mitochondrial functions. Electrophysiological changes in HD evolve over time in a regionally specific manner with circuit dysfunction encompassing both progressive and compensatory components [12]. Metabolic changes may follow a similar contextual and temporal progression. Mass action drives shifts in the balance of energy produced by glycolysis or oxidative phosphorylation on a temporal basis. Slowly accumulating oxidative damage may progressively inactivate some metabolic enzymes. Transcriptional dysregulation resulting in abnormal mitochondrial biogenesis and dynamics contribute to the need for compensatory responses. Accumulation of muh1 results in trafficking defects. Enzyme expression changes could result from homeostatic regulation and/or transcriptional dysregulation. Failure of the ETC may occur in CNS mitochondria late in the disease process but does not drive the transition from presymptomatic to symptomatic stages. This dynamic viewpoint can incorporate much of the reported experimental literature, as discussed below. The compilation and synthesis of published works presented here generates hypotheses to be tested in future investigations.

Fragments of muh1 translocate to the nucleus and bind to DNA, altering transcription factor binding and downstream gene expression [13]. This transcriptional dysregulation has been postulated to be both an early-stage event in HD pathogenesis and a later stage disruption of homeostatic compensation for mitochondrial dysregulation, endoplasmic reticulum and oxidative stress responses [14]. However, such homeostatic dysregulation could occur at the level of metabolic enzyme activities that operate on a more rapid and contextual basis than transcription. Indeed tissue specific variations in metabolic enzyme expression and activities support the specific energy demands of tissue specializations [15, 16]. Within the brain, regionally specific energy needs would be similarly expected to be supported by differential gene expression and activity levels, irrespective of muh1 effects.

The regional metabolic phenotypes associated with HD reviewed here occur on a heterogeneous background of energetic profiles that differ among excitatory and inhibitory neurons, the various glial cells and the multiple compartments within these cells [17]. Since the majority of HD mitochondrial studies focus upon regional enzymatic functionality, alterations at this more detailed level remain elusive and results represent the mix of mitochondria from all cells in the harvested samples. For explanations of the various experimental mouse models of HD, the reader is referred to recent reviews [18–20]. Briefly, mouse models vary from the rapidly progressive R6/2 which expresses the human exon I fragment with 120 or more polyglutamine repeats, to more slowly progressing knock in models such as the Q111, Q140 and Q175. None of the mouse models discussed appear to have phenotype until the polyglutamine repeats exceed 100 [18–20]. If not explicitly stated, all results reported below were made with respect to appropriate controls detailed in the original reports.
OVERVIEW OF ENERGY-GENERATING PATHWAYS

Energy metabolism must be viewed as a system of interacting biochemical pathways with the electron transport chain and oxidative phosphorylation as the final route to produce ATP (Figs. 1–5). ATP can be generated by glycolysis. Substrates other than glucose can feed into the TCA cycle pathways. Starting in the cytosol, glycolysis converts glucose into pyruvate, producing two ATP molecules per molecule of glucose. Pyruvate must then be transported across both the outer and inner mitochondrial membranes before being metabolized through the TCA cycle to produce the reducing equivalents NADH and FADH₂. This occurs through a series of enzymatic reactions initially on the cytosolic side of the outer mitochondrial membrane (OMM) creating fatty acyl-CoAs, which combine with carnitine to cross into the matrix via the acyl-carnitine/carnitine transporter. Once inside the mitochondria, fatty acyl-CoAs are oxidized through a cycle removing 2 carbon atoms at a time to generate NADH, FADH₂ and acetyl-CoA. Anaplerotic reactions degrade amino acids into TCA substrates. Glutamate and aspartate can be converted into α-ketoglutarate and oxaloacetate, both TCA intermediates, providing additional entry points for fuel into TCA. In addition acetyl-CoA, the normal substrate entering TCA, also can be diverted to produce cholesterol and fatty acids [23]. As demand for energy generating substrates increases, alternate fuels may be pulled into TCA from β-oxidation or glutamate oxidation.

Regulation of energy generation

Throughout all of these intersecting pathways, energy production is tightly regulated through localized pools, allosteric or substrate inhibition, and feedback pathways that tailor flux to current environmental conditions or demands. In the first step in glycolysis, the interaction of hexokinase with the voltage dependent anion channel (VDAC) in the OMM channels ATP exiting the mitochondria to directly fuel conversion of glucose into glucose-6-phosphate [24]. ATP itself allosterically inhibits phosphofructokinase [25], the second enzyme in glycolysis. The ratio of the concentrations of ATP and ADP ([ATP]/[ADP]) allosterically inhibits pyruvate dehydrogenase (PDH), as do the ratios [NADH]/[NAD⁺] and [acetyl-CoA]/[CoA]. In the TCA cycle, ATP inhibits isocitrate dehydrogenase (IDH); succinyl-CoA and NADH inhibits α-ketoglutarate dehydrogenase; and all of these plus citrate inhibit citrate synthase (CS). Conversely ADP or AMP activate PDH, CS and IDH in TCA and phosphofructokinase (PFK) in glycolysis. In PPP, NADP⁺ stimulates and NADPH can inhibit glucose-6-P dehydrogenase limiting glucose entry into this pathway [25]. Redox balance provides additional control on metabolic fluxes as oxidative changes in the NAD⁺/NADH ratio determines the pyruvate/lactate ratio through mass action and the direction of flux through PDH [25]. Cytosolic accumulation of NADH could also influence mitochondrial adenine nucleotide supply as NADH reduces VDAC permeability to ADP under physi-
Fig. 1. Alterations found in the human post-mortem HD striatum late in disease in the system of Intermediate Metabolism Pathways. For details and differences between caudate (Cau), putamen (Pu), globus pallidus (GP) and cortex, see the text. Intermediate Metabolism Pathways: Glycolysis, the principal catabolism of glucose to pyruvate is represented centrally. Branching off from glucose-6-P is glycogenolysis to the left and the Pentose Phosphate Pathway (PPP) to the right. In the mitochondria, pyruvate is converted to acetyl CoA which enters the TCA cycle, center bottom. Additional fuel can be generated from β-oxidation of fatty acids, which produces reducing equivalents, NADH and FADH2, and acetyl CoA. Glutamate metabolism, bottom right, can be derived from or provide fuel for TCA intermediates (anaplerosis). The Electron Transport Chain (ETC, very bottom) utilizes the NADH and FADH2 produced in all of these pathways to provide the driving force for converting ADP to ATP. Reactions specific to astrocytes (e.g. GS, PC and GDH) or GABAergic neurons (e.g. GAD, GDH) are included but not specifically separated. Enzymes and metabolites not discussed in this review are not labeled. Diagram adapted from [251]. [6–8, 45, 48–52, 70, 76, 77, 80, 81, 85, 125, 136, 137, 168, 170, 171]. **Note:** Bold indicates an enzyme expression or activity has been assayed. Bold black or red indicates upregulation. Bold grey or blue indicates downregulation. Bolding of an enzyme or metabolite name indicate protein expression or concentration alterations. Partial bolding or coloring indicates differential expression changes among subunits. Bold and/or coloring of arrows indicate activity changes. No change is indicated by a brown script font or a half open brown arrowhead. The default sans serif black font and thin black arrows with small heads indicate that no specific information is available. Combinations of symbol changes and asterisks (*) indicate conflicting reports from different laboratories.
Fig. 2. Changes to metabolic pathways determined from R6/2 striatum late in disease at 12 wk of age. Annotations as in Fig. 1 [53, 54, 56, 67, 68, 70, 73, 92, 104, 105, 110, 111, 113, 134, 136–138, 182, 204, 252, 253]. Note: Bold indicates an enzyme expression or activity has been assayed. Bold black or red indicates upregulation. Bold grey or blue indicates downregulation. Bolding of an enzyme or metabolite name indicate protein expression or concentration alterations. Partial bolding or coloring indicates differential expression changes among subunits. Bolding and/or coloring of arrows indicate activity changes. No change is indicated by a brown script font or a half open brown arrowhead. The default sans serif black font and thin black arrows with small heads indicate that no specific information is available. Combinations of symbol changes and asterisks (*) indicate conflicting reports from different laboratories.
Fig. 3. Changes to metabolic pathways determined from R6/2 brain at 2 wk of age. Annotations as in Fig. 1 [44]. **Note:** Bold indicates an enzyme expression or activity has been assayed. Bold black or red indicates upregulation. Bold grey or blue indicates downregulation. Bolding of an enzyme or metabolite name indicate protein expression or concentration alterations. Partial bolding or coloring indicates differential expression changes among subunits. Bolding and/or coloring of arrows indicate activity changes. No change is indicated by a brown script font or a half open brown arrowhead. The default sans serif black font and thin black arrows with small heads indicate that no specific information is available. Combinations of symbol changes and asterisks (*) indicate conflicting reports from different laboratories.
Fig. 4. Changes to metabolic pathways determined from STHdhQ111/111 cells compared to STHdhQ7/7 cells. Annotations as in Fig. 1 [63, 87, 94, 119, 120, 134, 204]. Note: Bold indicates an enzyme expression or activity has been assayed. Bold black or red indicates upregulation. Bold grey or blue indicates downregulation. Bolding of an enzyme or metabolite name indicate protein expression or concentration alterations. Partial bolding or coloring indicates differential expression changes among subunits. Bolding and/or coloring of arrows indicate activity changes. No change is indicated by a brown script font or a half open brown arrowhead. The default sans serif black font and thin black arrows with small heads indicate that no specific information is available. Combinations of symbol changes and asterisks (*) indicate conflicting reports from different laboratories.
Fig. 5. Transporters and transport pathways for energy substrate movement across plasma and mitochondrial membranes. Key transporters involved in moving metabolites into CNS cells and mitochondria. Changes indicated have been reported in different HD model systems. The malate-aspartate shuttle crosses the IMM on the left. The carnitine shuttle that brings FA from the cytosol into the matrix is on the right. ST represents numerous substrate transporters across the IMM and OMM for pyruvate, glutamate, malate, α-ketoglutarate, aspartate, citrate and succinate, some of which may be neuron or astrocyte specific. GLAST, GLT1 and GLUT1 are localized in astrocytes. EEAC1 and GLUT3 are localized to neurons. Annotations as in Fig. 1 [67, 73, 75, 85, 87, 104–107, 120, 133, 142, 143, 145, 163, 253, 254]. Note: Bold indicates an enzyme expression or activity has been assayed. Bold black or red indicates upregulation. Bold grey or blue indicates downregulation. Bolding of an enzyme or metabolite name indicate protein expression or concentration alterations. Partial bolding or coloring indicates differential expression changes among subunits. Bolding and/or coloring of arrows indicate activity changes. No change is indicated by a brown script font or a half open brown arrowhead. The default sans serif black font and thin black arrows with small heads indicate that no specific information is available. Combinations of symbol changes and asterisks (*) indicate conflicting reports from different laboratories.

ENERGY DEFICITS IN HD

Rather than talk about all mitochondrial functions lumped together, this review will focus specifically on which pathways are compromised in HD, with a focus on nervous tissue. Abnormalities in other tissues may be noted but will not be systematically explored as energy demand and regulation may be tissue specific [27]. For example, AMP kinase (AMPK), a sensor of energy status on both cellular and whole body scales, may be hypoactive in R6/2 muscle but hyperactive in brain tissue [27]. In addition, the cellular and subcellular compartments specializing in a particular pathway will be indicated. The prevalent idea that gradual mitochondrial dysfunction, through insufficiency or reactive oxygen species (ROS) production, is a driver of the aging process has recently
been called into question [28]. Similarly, the idea that less ETC activity may be harmful has also been challenged. In transgenic mice with compromised mitochondrial function lacking the complex I subunit Ndufs4, a lower oxygen environment triggering activation of hypoxia inducing factor (HIF-α) was neuroprotective [29]. In light of these reversals in thinking, a reanalysis of the mechanisms producing energy deficits in the HD pathogenesis may be in order.

**Longitudinal changes in energy metabolism in HD**

The idea of disease progression inherently contains the notion that the longitudinal path runs uni-directionally towards increasing dysfunction. However, homeostasis, the regulatory processes that maintain biological systems within reasonable operating boundaries, has a great deal of latitude for countering disruptions to produce proper function and well being. Concomitant expression levels of ion channels can vary greatly and still produce equivalent neuronal electrical activity [30]. Similarly, highly disparate circuit parameters can produce equivalent rhythmic network activity [30]. At each level of complexity, such systemic flexibility underlies homeostatic processes [30]. So when considering the number of enzymes throughout these energy generating pathways, restricted expression or activity of one enzyme would be expected to invoke compensatory shifts in expression levels for many others such that the system continues to provide sufficient energy for survival.

Metabolic changes in premanifest or prodromal HD stages may represent homeostatic response as opposed to signs of irreversible damage. In one cross sectional study of clinical blood biomarkers, the metabolites that were elevated at premanifest stages returned to normal levels during moderate HD [31]. In the DE5 mouse, which only expresses muh7 in medium spiny neurons, respiration utilizing complex II (CII) substrates was reduced presymptomatically but increased symptomatically [32], demonstrating stage-specific shifts in energy production. In the slowly progressing Q111 knock in mouse, increased rates of ATP utilization and reciprocal changes in brain PCr and Cr observed when muh7 is first detectable in the nucleus at 6 wk, were reversed by 13 wk, suggesting a homeostatic restoration occurred [33, 34]. Thus energy pathways appear to be homeostatically regulated during disease progression in both humans and HD mouse models.

A prime example of an energetic physiological response that can either promote disease at late stages or protect at early stages is that of AMPK. By end stage disease in the striatum and cortex of the most severe HD mouse model, R6/2 at 12 weeks, and in human postmortem HD brains, AMPKα1 appears to be translocated to the nucleus, where its increased activity leads to a decrease in the expression of the anti-apoptotic protein Bcl2, contributing to neuronal demise [35]. In contrast, early in disease progression, AMPKα1 may play a different role. The ratio of phosphorylated (activated) AMPK to total AMPK is greater than normal in 4 mo old Q111 mouse brains, one of the slowest progressing models of HD [36] (but see [37]). In mouse striatum expressing virally delivered muh7, a model where the initial development of toxicity can be followed, simultaneous overexpression of a constitutively active form of AMPK, AMPKα1GOF, reduces lesion size [38]. Similarly, enhancement of AMPK function through overexpression or metformin-induced activation of AMPK-mediated phosphorylation in both nucleus and cytosol are protective [38]. This latter result corroborates studies in cell lines. AMPKα1 mRNA and protein levels in STHdhQ111/1111 cells exceed that of ST/HdhQ7/7 cells suggesting that AMPK may play a role in maintaining the viability of the ST/HdhQ111/1111 cells. Low doses of metformin or overexpression of a constitutively active AMPKα1GOF protect ST/HdhQ111/1111 cells against serum-deprivation induced cell death. AMPKα1GOF also reduces soluble mutant htt levels. This neuroprotection is consistent with the general ability of AMPK to suppress ATP consuming processes and promote ATP production [39] although measures of mitochondrial size or function did not accompany these studies. For whole body regulation of glucose intake and energy production situated in the hypothalamus, AMPK activation decreases synthesis and increases oxidation of fatty acids leading to increased food uptake [40, 41]. When administered in the drinking water of R6/2 mice beginning at 4 wk, metformin reduced clasping time and extended lifespan (but not other measures) of males but not females [42]. Consistent with this, individuals with manifest HD motor symptoms in the ENROLL-HD study who were also taking metformin for diabetes had statistically significant, although quantitatively small, improvements in cognitive performance compared
to other HD patients and non-affected age-matched controls [43].

A longitudinal KEGG pathway analysis of altered R6/2 total brain protein expression neatly lays out the oscillating nature of protein expression changes over the course of this mouse’s life [44]. Three or more proteins were altered in glycolysis and OXPHOS at 2, 4, 8, and 12 wks, but not 6 wk of life [44]. Expression of TCA proteins was altered at 2 and 8 wk, while glutamate metabolism changed at 2 wk, PPP expression at 4 wk, and fatty acid metabolism at 8 wk [44](Figs. 2 and 3). Clearly compensatory or regulatory processes of energy metabolism are driving both very early and late disease progression-associated changes. Energy generation was upregulated early in life, in a manner suggesting that fluxes through the entire intermediate metabolism fluctuate in a contextual, stage specific manner [44]. Oscillations in the expression patterns of several enzymes suggest dynamic regulation at this level as well. For example, the expression of SDH subunit Ip is depressed at 2 wk, augmented at 4 wk and depressed again at 8 wk [44]. Fatty acid metabolism was implicated by mRNA changes at 6 and 9 wk [44]. The involvement of PPP, fatty acid and glutamate metabolisms in these changes suggest that a deficit in a single pathway or enzyme cannot explain the energy issues in HD; rather fluxes through the entire system must be considered.

Glucose uptake

Consistently, 2-[18F]-fluoro-2-deoxy-glucose (FDG) PET studies have documented progressively decreasing striatal and cortical glucose uptake in presymptomatic [45–47] and symptomatic individuals [45, 48–52]. PET FDG uptake documented decreases in glucose uptake in striatum and frontal and parietal cortices in patients symptomatic for <5 yr spread to additional areas as disease progressed further [50]. Depressed HD patients had lower glucose uptake in thalamus and orbital-frontal inferior prefrontal cortex [51]. In a longitudinal study of symptomatic patients, not only was the rate of decline in FDG uptake correlated with the severity of the UHDRS functional scores, but riluzole treatment was able to reverse both the motor and metabolic declines [52]. However, in presymptomatic HD carriers the story is more complicated. A metabolic network abnormality was identified that included decreased FDG uptake in striatum and anterior cingulate, plus increased FDG uptake in thalamus, vermis and primary motor and visual cortex [47]. Overall glucose uptake increased in this network 18 mo after the initial visit and then declined 26 mo later, suggesting early compensatory shifts in overall brain metabolism followed by decline once compensation was exhausted [47]. While glucose metabolism decreased continually throughout this study, thalamic metabolism first increased and then declined, driving the overall pattern [47].

Similarly, microPET or autoradiographic studies demonstrated reduced glucose uptake in the R6/2 mouse and an HD rat model [53–55] (but see [56, 57]). As in humans, impaired glucose uptake occurs early in the R6/2, with 2DG uptake being reduced in PFC, premotor cortex and striatum of R6/2 mice at 6 wk despite a simultaneously increased relative cerebral blood volume [53]. This may be another instance where metabolic homeostatic mechanisms attempt to compensate for changes induced by muhtt [53]. Notably, in the HD rat, FDG measurements were transiently elevated in sensorimotor cortex only at 2 months, otherwise no differences were observed in glucose uptake compared to controls [57].

Consistent with the PET studies, decreases in expression of glucose transporters (Fig. 5) have been widely reported in both humans and preclinical model systems. In postmortem HD brains at stage 3, but not stage 1, the expression of both the Glut1 (epithelial) and Glut3 (neuronal) glucose transporters is reduced [58]. Most importantly, increased expression of GLUT3 via copy number variations in humans delays the age of HD onset by three years, indicating the impact of boosting energy resources early in this disease [59]. Unfortunately, such copy number variations occur infrequently [59].

Limited glucose uptake, attributable to multiple mechanisms, may be one of the primary, initial restrictions on energy metabolism. Cell surface expression, but not total protein levels, of the GLUT3 glucose transporter is diminished in the striatum and cortex of homozygous Q140 mice compared to controls [60]. In 12 mo old BACHD mice, a decrease in glucose uptake was observed in the striatum, hippocampus and cerebellum pointing to a glycolytic deficit at a time when COX histochemistry, a functional ETC mitochondrial marker, did not reveal any differences attributable to muhtt [61]. Surprisingly, when undergoing whisker stimulation, cortical glucose uptake was greater in presumably presymptomatic BACHD mice at 64 wk than in the controls [61], suggesting a system whose responses are not well modulated. In primary co-cultures of astrocytes and neurons from BACHD mice, 2-deoxyglucose
uptake was diminished whenever muhtt expressing astrocytes were present, suggesting that a glycolytic deficit may be attributable to astrocytes rather than neurons [61]. Glucose uptake is attenuated in primary cultures from Q140 neonates as a consequence of elevated rab11 activity [62]. Glucose uptake is decreased in STHdhQ111/111 cells compared to STHdhQ7/7 cells due to decreased mRNA, protein and surface protein levels for the Glut3 transporter [63]. Glut3 mRNA is also reduced in R6/2 striatum as early as 3–4 weeks of age [63]. Glucose and its metabolite glucose-6-phosphate have been identified as stimulators of mTOR-induced autophagy, so a decrease in overall glucose uptake could also downregulate normal autophagy, resulting in more accumulation of muhtt aggregates aggravating disease [64]. Together, these studies suggest that altered Glut3 protein expression and trafficking contribute to the metabolic phenotype in HD.

**Glycolysis**

In brains of HD patients assayed with PET, the efficiency of conversion of glucose to ATP is maximal in a resting state [65]. Thus when brain activity increases, the ability of the energy generating biochemical pathways to respond rapidly with an increase in ATP production, does not occur or may be limited [66, 67]. Similarly in the occiput of affected individuals, onset of visual stimulation fails to increase P2/ATP, PCR/ATP or P/PCR ratios or lactate accumulation, as observed in control subjects [66]. The glycolytic enzyme phosphofructokinase was diminished in the post mortem HD caudate and putamen, but not in cortex [68] (Fig. 1). The PPP enzyme glucose-6-phosphate dehydrogenase, was unchanged, indicating that the decreased glycolytic flux was not caused by a diversion of glucose into the PPP [68]. Both normal and polyglutamine expanded muhtt bind to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) but GAPDH activity was not altered in post mortem human caudate, putamen, cortex or cerebellum [69, 70]. However, at 10 wk of age, iodoacetate inhibited GAPDH more in R6/2 striatum and cortex than in control brains, although no mechanism for this increased susceptibility was uncovered [71]. Differential proteomics of samples from stage 4 post mortem brain identified an increase of aldolase, as prominently elevated in both striatum and cortex [72]. Enolase was also carbonylated (oxidized), suggesting its activity would be compromised [72]. Flux through glycolysis may become limited by such protein oxidation as disease progresses. Both neuronal and non-neuronal forms of enolase, one of the later enzymes in the glycolytic pathway, become oxidized in R6/2 striatum at 10 wk, resulting in a lowered overall activity level [73]. Consistent with this, enolase protein levels decreased in both R6/2 cortex and striatum at a time when general protein carbonylation was increased [67]. Additional evidence that striatal neurons survive on glycolytic energy comes from the reduced level of cell death in R6/2 and R6/1 mouse brains induced by malonate, an inhibitor of succinate dehydrogenase, an enzyme that participates in both TCA and ETC [74]. This phenomenon occurs in older, but not younger R6/1 mice, suggesting that a shift (perhaps gradual) from oxidative metabolism to glycolysis occurs with disease progression [74].

While the abnormal polyglutamine expansion may produce a gain of toxic function, the absence of normal htt levels may contribute to a loss of normal function. In human embryonic stem cells (ESC) lacking huntingtin expression, dramatic shifts in energy metabolism, including glycolysis, are characterized by altered metabolite levels [75]. Compared to normal ESC, shifts characteristic of near maximal glycolytic flux include decreases of intracellular glucose, glucose-6-phosphate and fructose-1,6-diphosphate, increases in all glycolytic triose intermediates prior to pyruvate and increases in glucose uptake and pyruvate conversion to lactate [75]. In contrast, ESC expressing Q140/7 muhtt, pyruvate increases at the expense of lactate and many fatty acids and carnitines become overabundant [75]. Consistent with this, nucleotide triphosphate levels drop in htt/- cells and nucleotide di- and monophosphates increase in the htt/- cells [75]. Increases in metabolites involved in purine biosynthesis also appear to be part of a homeostatic shift in metabolic pathways to increase ATP synthesis [75]. Surprisingly mitochondrial membrane potential is maintained in both Q140/7 cells and htt/- cells, the latter presumably at the expense of glycolytically generated ATP [75]. It is noteworthy that the ETC was not altered in either htt/- or Q140/7 ESC [75]. While it is unclear how the metabolic adaptation to growing in cell culture may be influencing these results, the absence of normal htt function clearly shifts metabolism towards glycolysis and away from TCA and oxidative phosphorylation, not unlike the early phenotype described above in humans [65]. The presence of polyglutamine expanded muhtt also shifts metabolism, but less dramatically and less clearly in this model system [75].
**Tricarboxylic acid cycle**

The earliest studies of metabolic enzyme deficits in HD realized that activity might change with disease progression. In first assessments of HD postmortem brains, succinate dehydrogenase (SDH) decreased only in the most severe cases [76](Fig. 1). SDH was also depleted beyond what was expected from the agonal state in the caudate but not frontal cortex [77]. Aconitase activity was found to decrease in post mortem caudate, putamen and cortical areas BA 10 and 3, but not cerebellum [70, 72]. Differential proteomics of samples from stage 4 post mortem brain identified a decrease of aconitase and an increase in its oxidation in striatum, but not cortex, leading to a loss of activity in striatal samples [72]. Aconitase activity can be compromised by carboxylation (oxidation) or by aggregation [78, 79]. Tissue transglutaminase, whose activity doubles in grade 3 or 4 HD caudate due to muhtt binding, causes aconitase to aggregate within mitochondria [78, 79]. Similarly CS activity was decreased in putamen and cortex [70]. Activity of the PDH complex was decreased in the caudate, putamen and hippocampus, but not frontal cortex or amygdala of HD brains [80]. The correlation of the loss of caudate PDH activity with disease duration suggested progressive loss with increasing disease burden [77]. \(\alpha\)-Ketoglutarate dehydrogenase activity is also depressed in post mortem putamen [81].

In all analyses of post mortem human brain enzyme activities, experimenters have to take into account the effects of age, postmortem tissue interval and quality of tissue preservation. Agonal state, the physiological status of the patient prior to death and the suddenness of death, are rarely reported or considered. Slow deterioration in the last few weeks of life diminished respiratory rates of synaptosomes isolated posthumously from non-demented bronchopneumonia patients compared to those from sudden death cardiac arrest patients [82]. This was attributed to a decline in pH of cortex, CSF and blood presumed to arise from pre-mortem lactic acidosis in the agonal state [83]. In a broad study of mitochondrial DNA changes associated with depression in postmortem human brains, the biggest differences were observed in control brains where tissue pH predicted agonal state [84]. Few studies take into account the agonal state when reporting enzyme activities of SDH, PDH and aspartate amino transferase (AAT) [76, 77, 85]. Agonal state can also influence activity of glutathione reductase and glutathione-S-transferase [86]. While curatorial consistency has improved in contemporary brain banks [87–89], agonal state remains rarely considered.

The most recent metabolomic examination of TCA function in HD was undertaken with the intent of examining all TCA enzymes from the same material, to minimize differences in preparations, laboratories and physiological contexts. Naseri et al. examined the activities and mRNA and some protein levels of all TCA enzymes from postmortem HD parietal cortex and heterozygotic Q175 cortex and striatum at 14-15 mo [90, 91]. In the cortex from mostly grade 2 and 3 cases, an increase in SDH enzyme activity was reported [91]. A similar increase in SDH activity, accompanied by increases in PDH complex and aconitase were observed in Q175 cortex, but not striatum [91]. These results contrast sharply from the late stage post mortem profile, possibly reflecting regionally localized compensatory responses during mid-stage disease. In the Q175, the elevation of PDH activity at the entry to TCA bolsters the subsequent aconitase and SDH steps, representing an attempt to increase energetic flux. Strikingly, no changes were detected in the presumably symptomatic Q175 striatum.

TCA enzyme alterations in the rapidly progressive R6/2 mouse contrast to those in the slower Q175 model (Fig. 2). In R6/2 striatum at 10 wk compared to 4 wk, protein levels of dihydrolipoamide S-succinyltransferase, an E2 subunit of both the PDH and \(\alpha\)-ketoglutarate dehydrogenase complexes, increased but PDH protein decreased [73]. This has been interpreted to represent an attempt of one subunit to compensate for loss of activity of the whole complex [73]. Aconitase activity was also reduced in striatum but not cortex of R6/2 at 12 wk [92]. At 10 wk, aconitase protein from R6/2 striatum was more oxidized than at 4 wk, suggesting progressive inactivation, which would impede flux through TCA [67, 73]. By 13 wk, aconitase was oxidized in both regions [93].

The transcriptional dysregulation generated by muhtt may include increased expression of pyruvate dehydrogenase kinase (PDK), which acts to inhibit pyruvate dehydrogenase (PDH) activity [94]. Mechanistically, PDH activity is reciprocally down or up regulated when phosphorylated or dephosphorylated by isoforms of pyruvate dehydrogenase kinase (PDK) or pyruvate dehydrogenase phosphatase (PDP), respectively [95]. In ST\(HdhQ^{111/111}\) cells, observed decreases in PDH activity could be explained by increases in PDK protein levels and decreases in PDP leading to phosphorylation of...
PDH [94] (Fig. 4). Inhibitors of PDK or the HDAC inhibitor sodium butyrate ameliorated this phenotype and restored PDH activity, mitochondrial respiration and ATP levels [94]. A month long course of sodium butyrate also decreased PDK, phosphorylated PDH and restored motor behavior and relative ATP levels in the cortex of YAC128 mouse at 9 mo [94]. Why PDH activity is depressed in STHdhQ111/111 cells and possibly YAC128 cortex [94] but elevated in Q175 cortex [91] remains unclear. These differences may be attributable to the influence of the metabolic state of background strains or the extent of disease progression at the time the experiments were performed.

Anaplerosis and glutamate oxidation

In addition to generating energy, TCA substrates are used in the synthesis of the amino acids glutamate, glutamine, aspartate, and alanine in pathways which branch off of TCA, referred to as glutamate metabolism pathways. These reversible reactions can work in reverse to utilize amino acids as entry points into TCA, for example during glutamate oxidation by GDH. Alterations in these enzymes have also been reported in HD brains and models.

In postmortem HD globus pallidum, GDH was elevated, presumably because of its association with the abundant and increasing presence of astroglia in this brain area [68] (Fig. 1). GDH associates with mitochondria and glutamate transporters to efficiently shuttle exogenous glutamate into the TCA [96]. Within TCA, GDH forms multienzyme complexes with many enzymes to facilitate glutamate utilization as an energy substrate [96]. Their (dys)regulation could contribute to either pathogenic or compensatory shifts in energy production. Astrocyte glutamate synthesis derives from flux through pyruvate carboxylase, an enzyme only found in astrocytes, forming an anaplerotic pathway into TCA [97–99]. While flux analysis tracing carbon movement through this pathway [100] has not been done on HD models, pyruvate carboxylase protein levels were found to be elevated in cultured astrocytes from neonatal BACHD cortex [101], consistent with an increased flux through TCA and increased production of glial glutamate in presymptomatic brain.

Aspartate amino transferase (also known as glutamic oxaloacetic transaminase) reciprocally converts oxaloacetate and glutamate into aspartate and α-ketoglutarate. This is the entry point for glutamate conversion into energy via a partial TCA cycle [102, 103]. In R6/2 striatum at 10 wk, aspartate amino transferase expression increased, supporting the idea that flux through the metabolic networks shift in a compensatory manner [73] (Fig. 2). In the face of a reduced flux produced by aconitase oxidation and inactivation, this shortcut would provide an alternative path to supply α-ketoglutarate [73]. An increase in the use of glutamate to fuel this portion of TCA cycle occurs in rat brain synaptosomes under hypoglycemic conditions [102, 103]. While not as severe as the experimentally induced absence of glucose, the reduced glucose uptake in HD could invoke utilization of glutamate as an energy source. HD associated loss of AAT activity in postmortem caudate could indicate that availability of glutamate as an energy substrate was depleted by end stage disease [85] (Fig. 1).

Despite the reported decreases in protein expression for plasma membrane glutamate transporters in HD mice [104–106] (Fig. 5), functional glutamate uptake was normal or increased in YAC128 and R6/2 striatal slices on a synaptic activity time scale [107]. This rapid glutamate clearance could serve as a temporally tuned source of exogenous glutamate for energy generation. Indeed, association of astrocytic glutamate transporters GLT1 and GLAST with GDH and mitochondria, respectively, facilitate use of exogenous glutamate as an energy source [96], suggesting that increased glutamate uptake may be an adaptive anaplerotic response in HD brain.

Electron transport chain

As the set of common final reactions that produce the electrochemical gradient for driving ATP production, the ETC has been investigated as a locus for possible metabolic abnormalities in HD. In a large sample of post mortem grade 3 and 4 HD brains, CI and CII-III activity was depressed in caudate and putamen [6, 9] (Fig. 1). However, in presymptomatic or grade 1 striatum, activity of all the ETC complexes was normal [9], clearly indicating that major compromise of the ETC is a late event in pathogenesis. Largely overlooked in this study were sizable increases in Cl and CII-III activity in the cerebellum, a part of the brain comparatively resistant to muhtt [6]. Such shifts support the concept that metabolic disruption is a system–wide phenomena in HD that triggers regionally specific homeostatic compensations.

In mitochondria prepared from postmortem human HD brains, respiration and cytochrome oxidase activity were normal despite unknown post mortem intervals or disease stage at death [8]. A set of recent
In symptomatic, older N171-82Q mice CII or lactate/pyruvate fueled respiration decreased symptomatically [32]. Similarly, in young mice expressing the N171-82Q construct in striatum, CII, but not CI, fueled mitochondrial respiration only increased, conditions that were rescued by overexpression of the Fp or Ip subunits [125]. Significant differences were not detected in expression of Cox IV, Cyt C or the α subunit of CV [125]. In R6/2, expression of the Ip subunit oscillates over the course of disease [126]. In the R6/1 striatum at 16 wk, Ip subunit expression was reduced while in the N171-82Q mouse, CII expression was reduced beginning at 1

The apparently contradictory results obtained in mice expressing the N171-82Q construct must be viewed by considering that metabolic flux may shift over the course of disease. In young animals only expressing the N171-82Q construct in striatum, CII, but not CI, fueled mitochondrial respiration was reduced presymptomatically, and subsequently increased symptomatically [32]. Similarly, in young or presymptomatic N171-82Q transgenic mice, only CII or lactate/pyruvate fueled respiration decreased [114, 115]. In symptomatic, older N171-82Q mice an increase in CI activity [116] or a decrease in CI, CII or CIV activity have all been reported [114, 117, 118]. These results in the N171-82Q model suggest that metabolic shifts in end stage disease, while distinct from early stages, may be quite variable and dependent upon local husbandry or experimental conditions.

In STHdhQ111/111 cells, activities of all the mitochondrial complexes were equivalent to those from the non-polyglutamine expanded cells, yet mitochondrial respiration on CI and CII substrates (but not CIV substrates) was reduced in situ [119] (Fig. 4). This points to a failure of the entire ETC to produce adequate fluxes in the cellular context. In contrast, another examination of these same cells reported decreases in CI through CIV activities in homozygous cells, but only decreases in CI and CIV in heterozygous cells [120]. Altered expression of two mitochondrial disulfide relay system proteins, responsible for import of Fe-S containing proteins including many components of cytochrome oxidase, was offered as a mechanistic explanation [120]. Importantly, these changes were dependent upon gene dose and passage number [120], demonstrating the contextual nature of energetic deficits.

SDH, or CII, has been proposed as a target for muhtt-induced inhibition since exposure to low doses of chemical inhibitors over long periods mimics some HD phenotypes [121–124]. The fact that inhibition of energy pathways, especially of SDH, mimicked HD phenotypes motivated the search for underlying metabolic abnormalities. Contributing to both TCA and CII activity in ETC, SDH occupies a critical position metabolically. SDH activity, measured as CII, was depressed with respect to citrate synthase in post mortem caudate at presumably end stage disease [7]. In caudate and putamen, but not cortex or cerebellum of stage 1–3 HD brains, expression levels of the Fp and Ip subunits of SDH were depleted [125] (Fig. 1). Similarly, primary neurons transfected with Htt171-82Q had decreased expression of these subunits and consequently lower SDH activity. In addition mitochondrial membrane potential was depleted and TUNEL positive cells were increased, conditions that were rescued by overexpression of the Fp or Ip subunits [125]. Significant differences were not detected in expression of Cox IV, Cyt C or the α subunit of CV [125]. In R6/2, expression of the Ip subunit oscillates over the course of disease [126]. In the R6/1 striatum at 16 wk, Ip subunit expression was reduced while in the N171-82Q mouse, CII expression was reduced beginning at 1
Oxidative phosphorylation and ATP production

In humans with early stage HD, $^{31}$P magnetic resonance spectroscopy (MRS) of occipital cortex revealed normal basal ATP levels [66]. Only one study reported oxidation of CV in postmortem stage 4 striatum [72]. Variable deficits in basal ATP levels have been reported in preclinical HD models. In symptomatic R6/1 mice at 17 wk, decreases in cortical, but not striatal ATP were noted [130]. On the other hand, in Q150 knock in mice at 14 but not 4 mo, ATP levels were depressed only in synaptosomes, remaining normal in regional brain homogenates [131]. In 9 mo old YAC128 whole brain, ATP/ADP only tended to be decreased [94]. In symptomatic N171-82Q mice, an increase in ATP levels [116] and no change in cortical or striatal ATP/ADP ratios or striatal CV activity [32, 117] have all been reported. ATP was decreased in R6/2 striatum and cortex at 8 and 12 weeks [36]. However, $^{31}$P MRS of anesthetized R6/2 failed to find a decrease in the ATP peak [33]. In R6/2 mice at an unspecified age, ATP levels in striatum and cortex did not differ from controls, consistent with no changes in ETC activities, despite a reduction in mtDNA copy numbers [111]. While such differences could be methodological, a more likely explanation might be that the mice were more energetically stressed in one setting compared to another. Whether [36, 130, 131] or not [33, 131, 132] a laboratory finds decreased ATP levels in the HD brain may be dependent upon the physiological status of the animals and tissue at and following sacrifice, often overlooked variables. In human lymphoblast cell lines expressing normal and mutant htt, the ATP/ADP ratio decreased as the polyglutamine expansion lengthened [133]. Total ATP levels in the STHdhQ111/1111 cells have been reported to be decreased or the same as those found in their STHdhQ/77 counterparts [119, 120, 133]. Decreased CV activity was observed in STHdhQ1111/1111 cells, but not in STHdhQ111/7 cells [120]. CV activity was unchanged in primary neurons transfected with Htt171-82Q [125]. In Neuro2A cells expressing a muhtt exon 1, ATP levels were reduced [134]. In contrast, cultures of embryonic striatum from YAC72 mice exhibited normal ATP levels [132]. How ATP levels are altered in lymphoblasts, muscle, or immortalized cell lines may also depend upon the physiological context of those tissues [119, 120, 133, 135]. In vivo basal energy metabolism, through whatever pathway, may be adequate to supply resting state needs and inadequate during activation [66, 67, 131].

Cytosolic availability of ATP may be limited by muhtt-induced downregulation or oxidation of CK expression [72, 134, 136, 137]. Upon exiting the mitochondria, newly synthesized ATP is converted by CK to PCr, the cytoplasmic high energy storage molecule. Subsequently, cytosolic CK generates ATP from the PCr in response to local energy demands. Disruption of the conversion of PCr back to ATP at cytosolic sites where energy is needed could also restrict or disrupt cellular function and homeostasis. For example, proteostasis, a cellular function requiring local ATP that may be compromised in HD, was restored by CK overexpression [134]. Indeed, CK expression and activity becomes transcriptionally depressed in a longitudinal manner in patients and the R6/2 and Q140 mouse models of HD [134, 137] (Fig. 2). Single time point reductions in CK activity have also been demonstrated in older Q111 and N171-82Q mice [136]. In addition, CK becomes increasingly oxidized over the course of R6/2 disease [73]. As a result of CK depletion, PCr accumulates and local ATP may not be readily available.
energy source. In response to visual stimulation, 31P all restriction on the availability of the most abundant glucose uptake, discussed above, will place an over-

increased ATP production. Certainly the decline in ability to respond to an increased energy demand with the mitochondria within them must have the capac-

ity to bring substrate into cells and/or their mitochondria [141]. However, limited pyruvate, malate or NADH import could also explain these results.

Indeed, extramitochondrial calcium regulates glutamate fueled respiration by controlling aralar, the glutamate-aspartate transporter involved in the malate-aspartate shuttle, which imports NADH and glutamate and indirectly malate into mitochondria [142, 143] (Fig. 5). Thus, mitochondrial import of ETC substrates appear to be limited by cytosolic calcium. Additional restriction of substrate entry through the malate-aspartate shuttle might arise from an observed decrease in AAT activity in post mortem human caudate [85]. Substrate availability issues have surfaced in intact striatal neurons cultured from embryonic HD rat in which oxygen consumption rates, respiratory control ratio and spare capacity were normal in abundant glucose but decreased when glucose was restricted to 2.5 mM [144]. Alternate energy substrates (lactate and pyruvate) supplemented this respiratory function but did not fully restore it [144]. In STHdhQ111/Q111 cells, an observed decrease in ATP/ADP ratios was attributed to inadequate ADP uptake into mitochondria despite normal activity of the adenosine nucleotide transporter (ANT) [133], but restricted entry of upstream substrates could also have contributed. Comparing permeabilized STHdhQ111/Q111 and STHdhQ7/Q7 cells fueled by CI or CII, but not CIV, substrates, mitochondrial state 3 respiration, respiratory control ratios and the rate of ATP production were decreased with the polyglutamine expansion [119]. Notably under these conditions STHdhQ111/Q111 cells maintained normal ETC enzyme activity levels and sensitivity to inhibitors of complexes I through IV and normal ATP levels [119] (Fig. 4). The ETC machinery functioned normally but energetic flux appeared to become limited by some unidentified factor [119]. Given that plasma membranes were permeabilized, the limiting factor(s) must reside at the level of substrate transport across either the outer or inner mitochondrial membrane or a paucity of NADH generated by TCA. The pyruvate and phosphate carriers exert reasonably high controls on respiration in isolated brain mitochondria [145] but their function has not been explored in HD models.

Fatty acid β-oxidation

β-oxidation utilizes fatty acids to provide an alternative fuel source for energy production in the brain.
Somewhat contradictory evidence suggests that fatty acids may be utilized in the brains or bodies of individuals with HD. Metabolomic studies of symptomatic, but not asymptomatic, HD gene carriers’ plasma under controlled dietary and medicinal conditions revealed lower plasma levels of branched chain amino acids, carnitines, phosphatidylethanolamines and increased levels of fatty acid breakdown products suggesting that both β-oxidation and amino acids were being shunted into TCA as energy substrates [146]. Mass spectrometry analysis of frontal lobe and striatum from postmortem HD brains identified but-2-enioic acid, an intermediate in fatty acid biosynthesis, as elevated in cortex but depressed in striatum, another indication that β-oxidation may be utilized in a regionally specific fashion [147]. In this same study, metabolites in a number of amino acid synthetic pathways were also reported to be largely downregulated [147]. Analysis of human serum from presymptomatic gene carriers contained some of the same plasma catabolic markers but did not reach statistical significance [148]. In early stage HD patients, one month of dietary consumption of triheptanoin, a triglyceride with three 7-carbon fatty acids, improved the occipital Pi/PCR signal and increased levels of fatty acid breakdown products suggesting that both β-oxidation and amino acids were being shunted into TCA as energy substrates [148]. In this study, metabolites in a number of amino acid synthetic pathways were also reported to be largely downregulated [147]. Analysis of human serum from presymptomatic gene carriers contained some of the same plasma catabolic markers but did not reach statistical significance [148]. In early stage HD patients, one month of dietary consumption of triheptanoin, a triglyceride with three 7-carbon fatty acid chains, improved the occipital P/PCR signal detected noninvasively by 31P-MRS in response to visual stimulation [149]. Triheptanoin is metabolized in the liver via FAO to C5 ketone bodies, propionyl-CoA and acetyl-CoA, which can cross the blood brain barrier and serve as fuel sources [149]. This is a dramatic demonstration of the importance of alternative fuel sources for enhancing brain metabolism in HD.

β-oxidation and a pro-catabolic phenotype were implicated in an unbiased principle component analysis of serum metabolites discriminating between presymptomatic (8 wk) and symptomatic (15 wk) N171-82Q mice [148]. However, in N171-82Q mouse serum, mRNA levels of the intermediate enzyme in the β-oxidation pathway, acyl-coenzyme A dehydrogenase, are significantly lower at 13 wk but not at wk compared to controls [115, 117]. L-carnitine, a cofactor required for transport of fatty acids into mitochondria in the rate limiting step in β-oxidation, may also ameliorate symptoms in N171-82Q HD mice after long term IP injections [150] (Fig. 5). N171-82Q mice receiving IP injections of L-carnitine had an extended lifespan, fewer nuclear muhtt aggregates, reduced neuronal loss and improved motor performance [150]. However, whether this protection stemmed from facilitation of β-oxidation or antioxidant properties of L-carnitine was not established.

Cytosolic fatty acid synthesis and mitochondrial β-oxidation are normally mutually antagonistic cellular processes. Surprisingly, evidence for possible activation of both β-oxidation and FAS stems from a recent unbiased multiomics analysis of the STHdhQ111/111 and STHdhQ7/7 cell lines [151]. This study iteratively matched partially identified differentially expressed mass spec peaks between the two cell lines with an integrated data base of known metabolic pathways, lipids and proteins. This process produced networks of connected, up- or down-regulated metabolites and inferred abnormalities in the biochemical pathways that connected them. Changes in metabolites that were predicted to be abnormally expressed at intermediate positions on the identified pathways were subsequently validated by conventional methods. From the protein fragments in the original mass spec analysis, abnormalities in sphingolipid, fatty acid and steroid metabolism emerged. This includes a demonstration that protein expression of FAS and two predicted long chain fatty acids (EPA & DGHLA) were elevated in STHdhQ111/111 cells compared to STHdhQ7/7 cells [151] (Fig. 4). FAS, responsible for de novo fatty acid synthesis downstream of acetyl-coA, also regulates food intake in the hypothalamus [152]. EPA stabilizes mitochondrial integrity and increases β-oxidation [23]. The prominent alterations in fatty acid metabolism and upregulation of these competing pathways suggests that the STHdhQ111/111 cell line may utilize so much fatty acid as fuel that its general synthesis needs upregulating as well. If these pathways are also increased in human brain, this may be another example of a homeostatic response to muhtt.

Compensatory shifts to generate energy from fatty acids or amino acids may be regulated by sirtuins, which deacetylate enzymes in fatty acid oxidation and amino acid metabolism [153]. SIRT3, whose short form is located in the mitochondria, becomes co-activated by NAD+ to deacetylase enzymes involved in fatty acid oxidation, promoting flux through this energy-generating pathway [153]. NAD+ levels are rate limiting for SIRT3 deacetylation, thus SIRT3 becomes a sensor of metabolic flux. In peripheral tissues, SIRT3 levels increase during caloric restriction or periods of lowered glucose availability and decrease with aging [153]. SIRT3 forms part of a positive feedback pathway to increase energy metabolism through activation of AMPK by increases in AMP levels. AMPK activated CREB and subsequently PCG-1α stimulate SIRT3, which in turn deacetylates LKB1, further stimulating AMPK [153]. SIRT3 also
deacetylates both isocitrate dehydrogenase 2, a TCA cycle enzyme producing NADPH, and Mn superoxide dismutase, a mitochondrial antioxidant enzyme, implicating SIRT3 in control of further metabolic flux and antioxidant defenses. During metabolic stress or low glucose availability, SIRT3 deacetylates a long list of mitochondrial proteins involved in TCA, β-oxidation, ETC and oxidative phosphorylation, all in the direction of promoting energy generation [159]. SIRT3 knock out mice are more vulnerable to oxidative stress, while running on a wheel increases SIRT3 expression, producing neuroprotection [154].

However, SIRT3 deacetylation may not be happening in HD due to sirtuin deficiencies, as observed in several HD models [37]. Indeed, muhtt inhibits general cellular deacetylase activity by binding to SIRT1 [155]. Binding of muhtt to SIRT3 has not been shown. Sirt1 overexpression improved functional measures in the N171-82Q mouse, suggesting loss of its regulation may influence pathogenesis [155]. However, Sirt1 levels at different stages of disease were not determined, so at what stage SIRT1 deregulation contributes to pathogenesis remains a mystery. In the ST/HdhQ111/111 cell model (compared to ST/HdhQ7/7 control cells), muhtt produces decreases in SIRT3 expression, pAMPK relative to AMPK and NAD+/NADH while increasing acetylated LKB1 and ROS levels [37]. These changes could be reversed by viniferin, a natural analog of resveratrol, an activator of SIRT3 [37]. Thus in these immortalized cells, muhtt expression suppresses the sirtuin pathway involved in compensating for energy deficits. However SIRT3 involvement in regulating fatty acid oxidation and acetylation status of specific enzymes remains to be explored. Furthermore, if muhtt decreases SIRT3 and subsequently decreases the suprarregulatory AMPK pathways involving both SIRT3 and PGC1α, this points to an eventual loss of master regulatory control over energy generation, consistent with the severe phenotype encountered at end stage human HD.

**Pentose phosphate pathway**

Involvement of increased fluxes through the PPP in HD is currently supported only indirectly. In the cortex of heterozygote Q175 mice at 14-15 mo, mRNA levels for transketolase, a rate-limiting step in the PPP, was elevated [156]. Activity levels were not determined [156]. Additional arguments for PPP involvement invoke the need to produce NADPH for compensation for oxidative damage. In samples from postmortem stage 4 HD brains, expression of the antioxidant proteins peroxiredoxins, glutathione peroxidases, SOD and catalase were increased in striatum and cortex [72]. At 2 wk but not later, transketolase expression was upregulated in R6/2 striatum [44] (Fig. 3). These suggest an increased antioxidant need and possible upregulation of the PPP, which produces NADPH for GSH regeneration from GSSG.

In the Drosophila eye model of HD, increased flux through the PPP induced by downregulation of ribose-5-phosphate isomerase was protective, demonstrating that PPP activation could be beneficial in HD [157]. Oxidative stress increases demands for reducing equivalents NADPH and NADH to regenerate the antioxidants thioredoxins and glutathione. Depletion of NADPH pulls glucose through the PPP. Indeed, in a drosophila model of HD, overexpression of the human neuronal glucose transporter (GLUT3) or glucose-6-phosphate dehydrogenase (G6PDH, the enzyme pulling glucose into the PPP) but not phosphofructose kinase (PFK, the second enzyme in glycolysis) extends the muhtt fly lifespan [158]. Moreover, overexpression of all three ameliorate the severe curtailment of lifespan induced by knockout of a subunit of pyruvate dehydrogenase (PDH) or a subunit of CI [158]. This latter observation demonstrates very clearly how endogenous shifts in metabolic flux might compensate for defects in later stages of energy production. A similar mechanism may be at work in human HD as a compensatory homeostatic response, as increased copy number for the SLC2A3 gene encoding GLUT3 increases expression of GLUT3 and correlates with increased age of onset [59]. While these fly studies raise possibilities, upregulation of the PPP in HD has not yet been systematically examined.

In line with this, neonatal (but not adult) iron supplementation in R6/2 mice increases oxidative stress, resulting in accelerated disease progression [159]. The rise in striatal and cortical GSSG at 12 wk associated with iron supplementation since weaning could arise because the PPP failed to compensate. A somewhat transient rise in GSH was found in normally raised R6/2 cortex at 8 wk, as if the system were adjusting homeostatically to attempt to compensate for an increased oxidative load [138]. Consistent with this, a rise in lactate accumulation in cortex of the R6/2 mouse suggests an increase in glucose utilization (beginning at 8 wk, significant by 12 wk) [138], perhaps including augmented flux through the PPP. Comparable changes in lactate concentrations in other models or human studies have not been reported as brain MRS lactate quantification often exceeds the
accepted variability limits [160]. Protein oxidation in R6/2 cortex was detected at 10 wk and beyond [67, 161].

The increased oxidative stress in HD may not come solely from an increased production of ROS by mitochondria. Rather the apparent increased oxidative damage may result from a decrease in antioxidant defenses unable to handle the normally generated level of spurious oxidation. In a metabolic environment with a restricted glucose supply, ATP generation may be favored over the PPP production of NADPH. Glutathione synthesis may be limited by mitochondrial or the huntingtin-associated protein-1 may be responsible for the dearth of plasma membrane SVCT2 transporters, contributing to a decrease in intracellular ascorbate, an increase in ROS levels and indirectly promoting glucose uptake over lactate uptake [63, 163].

The full extent of oxidative damage in HD is beyond the scope of this review. The reader is directed to discussions by other authors [161, 164].

Metabolite changes

Metabolomic analysis of steady state levels of small molecules in the brain relevant to energy generation corroborate a view of shifting adjustments at different stages of disease. Changes early in life or presymptomatically may represent homeostatic ones, whereas progressive changes may represent developing dysregulation. Uneven rates of change in metabolite levels were first reported in a longitudinal magnetic resonance spectroscopy (MRS) study of cortex and striatum from R6/2 [138] (Fig. 2). While choline-containing compounds and glutamine increased steadily from 4 to 16 weeks in both regions, elevations in creatine and phosphocreatine jumped initially between 4 and 8 wks and then remained largely stable [138]. N-acetyl aspartate (NAA) decreased beginning at 8 wk but more so in cortex than in striatum [138]. Myo-inositol increased earlier in striatum than cortex, while lactate increased more in cortex, suggesting that metabolic shifts might progress at rates influenced by regional connectivity and composition [138]. Somewhat similar but not identical shifts occur in the slower developing knock in mouse models. At 12 mo in the striatum of Q140 mice, glutamine increased while choline-containing compounds, glutamate, total NAA and taurine decreased [165]. The loss of glutamate was confirmed with chemical exchange saturation transfer imaging and further detected in cortex and corpus callosum as well [165]. In striatum of zQ175 knockout mice by 12 mo increases in glutamine, taurine, and total creatine, and decreases in NAA were reported [166]. Most interestingly, decreases in glutamate and GABA observed at 4 and 8 mo recovered by 12 mo, another illustration of a homeostatic compensation [166]. An additional study of the zQ175 striatal metabolites reported glutamate decreased at 6 mo, recovered at 9 mo only to decrease again at 12 mo, suggesting a regulatory system that is oscillating as disease progresses [167]. In addition, this study detected decreased NAA from 6 mo, increased glutamine from 9 mo and decreased GABA, and increased myo-inositol and taurine at 12 mo [167]. The differences in the latter two studies suggest that rearing conditions in different laboratories may also contribute variability.

Human MRS metabolite studies are more difficult to compare, as many studies report ratios of metabolite concentrations rather than the absolute values of concentrations. Fewer metabolites are typically detected due to use of lower field strength magnets. Consistent with the observation of early changes in R6/2 cortex, in premanifest to early HD, reduced NAA and glutamate levels in posterior cingulate cortex correlated with a measure of mild cognitive decline [160]. In the putamen, premanifest HD was characterized by a loss of NAA that continued to decline into early HD along with the development of a decrease in total creatine and glutamate and an increase in choline-containing compounds and myo-inositol [168]. The loss of NAA correlated with the total disease burden score and a number of
specific motor tasks [168]. The ratio of myo-inositol to NAA correlated with the UHDRS motor score [168]. In a 2 yr follow up longitudinal study, these changes persisted but did not progress making the correlations quite variable [169]. The loss of NAA and creatine have been confirmed by other laboratories [170, 171]. Proton MRS of HD patient CSF detected decreases in both lactate and citrate, but not in glucose, pyruvate, glutamine, alanine, creatine, creatinine, inositol, formate or acetate [172]. The changes in NAA and myo-inositol seen in humans parallel those reported in mice. However, the human MRS measurements remain quite variable and do not extend over a long enough span of disease progression to distinguish compensatory from degenerative changes.

Flux analysis

To truly capture the shifts in energy generating capacities, experiments must move beyond steady state determination of individual metabolite concentrations. Metabolic flux analysis and its derivatives provide a systems level view of reaction rates in enzyme pathways at metabolic steady state [173] and the ability to compare these fluxes across disease progression. Flux analysis applied to E Coli revealed that when and where carbon sources enter a metabolic network determine the overall energetic costs [174]. This suggests that given chronic high energy demands, cells may switch their reliance upon specific energetic pathways attempting to optimize ATP production. Comparing differentiated astrocytes and neural stem cells with \(^{13}\)C flux analysis after differentiation from embryonic stem cells, only astrocytes downregulated their carbon utilization [175]. Given that metabolic rerouting occurs during developmentally appropriate growth, similar metabolic adaptations could be expected to occur upon introduction of abnormal genotypes. With mhtt ubiquitously expressed from conception, tissue-specific metabolic adaptations should be tailored to each developmental stage. In liver glucose metabolism, the range of metabolic regulatory compensation achieved by rapid enzyme phosphorylation or allosteric or hormonal control of enzyme activities was equal to or greater than regulation by longer term alterations in enzyme expression levels [176]. For cerebellar granule neurons maintained in high glucose cultures, flux analysis demonstrated a high proportion of glucose was processed through the pentose phosphate and anaerobic glycolysis pathways [177]. Evolution of flux analysis techniques to in vivo conditions in the context of neuronal-glial interactions and normal electrophysiology, holds promise that the details of how metabolic networks shift in response to disease can indeed be worked out. However, caution must be exercised in identifying initial assumptions, scope and regional specificity when applying these broad systemic analyses [178].

Only one study has tried to look at metabolic shifts in HD from glycolysis through oxidative phosphorylation [116]. Olah et al. [116] went beyond measuring individual enzyme activities and modeled expected changes in fluxes, stating “the flux is a collective property of all the enzymes in the pathway.” In affected parts of the brains of 20 wk old N171-82Q mice, they reported the non-intuitive result that despite a decrease in GAPDH activity, an intermediate glycolytic enzyme, glycolytic flux increased as demonstrated by increases in pyruvate and lactate levels and activities of pyruvate kinase, enolase and hexokinase. The observed direction of changes in these products and enzymes agreed with predictions of their kinetic model [116]. Additionally, this study reported an increase in glutamate dehydrogenase (GDH) activity, an anaplerotic reaction which converts glutamate to \(\alpha\)-ketoglutarate, bringing this substrate as fuel into the TCA cycle, bypassing glycolysis [116].

In CNS mitochondria, the rate of ATP production is limited equally by CV and both the phosphate and pyruvate carriers, followed closely by CI [145, 179]. So inhibition of flux at any of these control points could compromise oxidative phosphorylation [145]. Examination of the control coefficients for metabolic flux compared across tissues reveals that brain is more sensitive to compromise of ATP synthase and the phosphate & pyruvate carriers than muscle and heart [145]. Liver is a close second in sensitivity to CV and the carriers, but is more sensitive to CI than brain [145]. Most notably, this study pointed out that the substrate carriers, which are often overlooked, are equally important when considering limitations on energetic flux [145]. A defect in substrate transport could account for the reduced respiratory control ratios in the absence of ETC enzyme deficiencies in STHdhQ111/111 cells discussed earlier [119]. Defects in substrate availability, due to a decreased VDAC1 expression, were also suggested to underlie the reduction in maximal in vivo cerebral metabolic rate of oxygen consumption (CMRO2) reported for R6/2 striatum and cortex at 10 wk [67].
Analysis of flux control ratios for each of the ETC complexes in R6/2 striatal homogenates at 12 wk uncovered significant but small decreases in CII and CIV control, with decreases in CII exerting more effect than CIV [180]. However, the overall respiratory control ratio was unchanged indicating that basal ETC capacity was normal and CII became the potential bottle neck if it became inhibited [180]. Endogenous inhibition of forebrain SDH occurs due to oxaloacetate binding [181], but this has never been examined in HD. Determination of control coefficients are rarely performed for other aspects of metabolic flux, such as glucose uptake, glycolytic enzymes or other substrate transport. The role of substrate flux, through the pyruvate carrier or the malate-aspartate or glycerol-3-phosphate shuttles that bring in NADH and FADH2, is an overlooked aspect of metabolic function in HD.

Ischemia reperfusion injury provides an informative example of how flux through all these energy generating pathways can shift with changing conditions. In heart, brain, liver and kidney, succinate becomes elevated during reperfusion [129]. In ischemia, abundant AMP drives its breakdown via the purine nucleotide cycle, producing fumarate. Fumarate however does not accumulate as it can be rapidly converted to succinate via the reversible SDH reaction. Subsequent tracing of labeled carbons, metabolic modeling and manipulation of succinate levels and SDH activity determined that during ischemia, fumarate converts to succinate reversing CII, consuming electrons from CI that were not pulled further through complex III (CIII) and CIV in the absence of oxygen. Upon reperfusion, the accumulated succinate drives normally directed flux through CII, producing excess electron flow that splits between the normal direction to CIII and CIV and reverse flow through CI generating reactive oxygen species [129]. As succinate levels decline, ETC flux returns fully to normal. Mass action shifts in flux through TCA cycle demonstrated by a decrease in aconitase activity, the alanine amino transferase (ALAT) and AAT reactions and the malate-aspartate shuttle during ischemia also would return to normal directionality as succinate concentrations subside [129]. Such short term temporal variations in metabolites have not been studied in HD model systems. Succinate was observed to decrease in R6/2 cortical and cerebellar, but not striatal, tissue extracts at 12 wk [182]. In presymptomatic HD rats, serum elevation of succinic acid was attributed to SDH inhibition [183].

In acute ischemic injury, expression levels were not altered, only fluxes. In HD, altered expression levels, in addition to mass action effects, will contribute to changes in flux that may be more complex than in ischemic injury. However, the mass action effects, as illustrated in ischemia, have explanatory power that must not be overlooked.

Classical concepts in metabolic regulation include the idea that energy demand drives production. However, recent flux analysis of metabolites and oxygen consumption in retina determined that phototransduction, possibly less energy demanding than maintaining ionic gradients for depolarization in the dark, slowed energy production through glycolysis and less so through TCA cycle [184]. Notably, mouse retinal oxygen consumption after light adaptation operated very near maximum uncoupled rates [184]. Flux analysis of this mostly rod-driven phenomena revealed that retinas relied much more on glycolysis in the dark. In the light, the PPP was stimulated and flux through TCA was slightly slowed [184]. The latter was accomplished through the light-induced decreases in free calcium which down regulated the malate-aspartate shuttle and the activity of α-ketoglutarate dehydrogenase [184]. Thus the metabolic consequences of changing energy demands may not be readily predictable. Similarly unpredictable, visual stimulation of occipital cortex in HD patients did not produce the elevation in Pi/ATP observed in normal subjects [66]. In anesthetized R6/2 in both cortex and striatum, the maximum uncoupled CMRO2 was less than in wild type mice [67]. Both of these experiments suggest that brain metabolism in HD may operate near maximum. While conditions in the HD brain are unlikely to exactly match those in retina, these nuances in retinal metabolism are examples of temporal and contextual energy regulation.

In cancer cells and immortalized cell lines, flux through metabolic pathways shifts to favor aerobic glycolysis over oxidative phosphorylation, a process called the Warburg effect [185, 186]. Induction of anaerobic glycolysis promotes cell longevity, enhancing tumorigenesis [186]. Such shifts have been found in response to synaptic activity in cultured hippocampal neurons on the fairly rapid time scale of hours [187]. Indeed, synaptic activity induced increases in gene expression for glucose and lactate transporters and PDH but a decrease in PDH activity within a few hours [187]. In the AD brain, the Warburg effect coincides with the development of resistance to Aβ accumulation and toxicity on a longer time
scale, where resistance was characterized by upregulation of LDHA and PDK favoring lactate production and decreased entry into TCA, respectively [188, 189]. Also associated with the shift to glycolysis and downregulation of oxidative phosphorylation was a decrease in potentially damaging ROS [188, 189]. Hypoxia inducible factor 1 (HIF-1) has been designated as a switch capable of upregulating glycolysis by induction of LDHA and downregulating TCA through reduction of PDK [190]. These AD and cancer studies present two ideas that must be considered in analyzing metabolic shifts in HD. The first is that the immortalized or cultured cells used in many HD studies may already have shifted set points favoring glycolytic activity. To their credit, investigators utilizing such cells compare the polyglutamine expansion to non-expanded cells. The second and more important idea is that the Warburg effect must be considered as a potential baseline condition in HD brain, where surviving neurons generate energy largely by glycolysis. Indeed HIF-1 induced decreases in PDK have been demonstrated in STHdhQ111 cells and YAC128 cortex [94].

Urea cycle

Difficulties in buffering brain amines may be an overlooked phenotype of brain metabolism in HD. While the brain contains urea cycle enzymes, normally this pathway is not thought to play a major functional role in nitrogen fixation [191]. Ammonia fixation in the brain largely occurs via amination of glutamate to glutamine by glutamine synthase in astrocytes [99, 192]. Not only was glutamine progressively elevated in the R6/2 striatum and cortex [138, 182] (Fig. 1), but 31P MRS of human brain revealed an increased intracellular pH [193]. Such alkalinization could reflect abnormalities in ammonia fixation in brain or liver or both. Metabolomic analysis of postmortem HD brain samples revealed an elevation of urea uniformly across eleven brain regions, including ones not affected by HD [194]. Consistent with this rise, the urea transporter UT-B present in astrocytes is elevated in postmortem human caudate and cortex, possibly to facilitate removal of the excess urea [194, 195]. A deficit in urea cycle functioning was noted in a prior report demonstrating decreases in postmortem striatal urea indicating additional work is needed to examine enzyme activities and to clarify how the urea cycle may be altered in HD [196].

While often overlooked, pH is a powerful regulator of enzyme activity. For example, phosphofructokinase is stimulated by ammonia while α-ketoglutarate dehydrogenase is inhibited, combining to shift energy flux from oxidative phosphorylation to glycolysis [197]. In the liver, the urea cycle detoxifies ammonia, preventing its accumulation in blood and subsequent widespread alkalinization of body tissues, including brain. Increased levels of ammonia and/or the urea cycle intermediates citrulline and arginine in blood from HD patients, HD sheep, and R6/2 and Q150 mice all point to abnormalities in liver urea cycle function [198, 199]. Indeed, lowering ammonia production with a low protein diet begun before symptom onset ameliorated accumulation of aggregates in the liver and striatum, partially restored liver mRNA levels of a transcription factor for urea cycle enzymes, ameliorated changes in activity of these enzymes, improved motor performance and restored striatal BDNF levels in R6/2 mice [198]. If indeed these peripheral effects of urea cycle abnormalities are early signs of HD onset, as illustrated in the plasma of presymptomatic HD sheep [199], the CNS metabolic defects could reflect a consequence of abnormal pH regulating various metabolic enzymes.

Changes at the organelle level

Mitochondrial biogenesis

The metabolic consequences of decreased PGC1α, a master regulator of mitochondrial biogenesis, have become well established in HD pathogenesis [2, 115, 200]. PGC-1α acts as a co-regulator in conjunction with the nuclear transcription factor PPAR-γ to regulate many genes involved in mitochondrial formation [115]. In humans, two distinct single nucleotide polymorphisms in the PPARGC1A gene modify the age of HD onset in opposite directions [201]. Muht interferes with PCG-1α striatal transcription, negatively impacting mitochondrial function [2]. PGC1α mRNA levels were decreased in samples from postmortem human HD striatum as well as in several, but not all, mouse and cell models (Q140, N171-82Q, STHdhQ111, but variably or not in R6/2) [2, 115, 202, 203]. Mitochondriobiogenesis decreased in PC12 cells expressing muht [204]. In the human samples, compensatory upregulation of mRNA for PPAR-γ and other transcription factors downstream of PGC-1α were also detected (64). PGC-1α overexpression in N171-82Q mice increases expression of mitochondrial enzymes, ameliorated motor phenotype and oxidative stress and prevented...
aggregate formation and neurodegeneration [117]. Similarly, overexpression of PGC1α restores mitochondrial function and prevents cell death among STHdhQ111/Q111 cells [2]. Rosigliten, a PPAR-γ agonist, rescues striatal cells from muhtt toxicity and restores motor function, BDNF levels and PCG1α levels and prevents hypothalamic neuronal loss in the N171-82 mouse model of HD [205]. Stimulation of PPARs with dietary thiazolidinedione or bezafibrate improved mRNA expression of mitochondrial enzymes in brains of R6/2 mice, increased striatal mitochondrial density and ameliorated neuropsychopathology and behavior [206, 207]. Restoration of mTORC1, an upstream positive regulator of PGC1α, also increases expression of PGC1α and associated mitochondrial enzymes and counteracts abnormal behaviors, again in N171-82Q mice [208]. In many of the overexpression rescue experiments, PGC1α and associated mitochondrial enzymes were elevated to greater than normal levels, suggesting that given enough energy, cells can overcome the proteostatic burden of accumulating muhtt.

Despite this strong genetic link to mitochondrial biogenesis, PGC1α knock out mice are viable [202, 209]. In normal development, PGC1α expression colocalizes with and peaks in striatal, cortical and hippocampal GABAergic neurons during the first several weeks postnatally in mouse [210]. In the PGC1α knock out mouse, an early onset but not progressive hyperactive phenotype and claspings are associated with striatal vacuolization and possible white matter loss [202, 209]. These mice differ from R6/2 HD mice in that markers for medium spinal neurons increase in a homeostatic manner, supporting the idea that metabolic pathways do shift to maintain function [202]. Given that in HD, PGC1α and the linked metabolic protein transcription may only be partially depleted [2, 115, 202, 203, 211], compensatory mechanisms may also be present. More importantly, PGC1α appears to more profoundly regulate expression of parvalbumin (PV) and several synaptic proteins critical for transmitter release to a greater extent than mitochondrial proteins [211, 212]. While the transcriptional mechanism for PGC1α suppression of PV has not yet been identified, such regulation suggests PGC1α depletion may cause dysfunction through non-metabolic means [211, 212]. Indeed, mice expressing muhtt exon one in PV expressing neurons exhibit an HD-like hyperactive motor phenotype and age-related changes in inhibitory synaptic events that may be independent of transcriptional dysregulation, suggesting that muhtt and PGC1α driven changes in PV neurons may also be independent of effects upon mitochondrial proteins [213].

**Mitochondrial trafficking**

The most energy-consuming communication processes of the nervous system occur at synaptic sites, far from the soma. To deliver energy to remote processes, mitochondria must be trafficked to dendrites and axon terminals, a process that can be disrupted by mutant huntingtin [4]. N-terminal mutant huntingtin impairs mitochondrial trafficking in axons of cultured striatal neurons from Q150 knock in and YAC72 mice and primary rat neuronal cultures transfected with muhtt [131, 132, 214]. Both anterograde and retrograde transport were slowed [131, 132]. Full length muhtt slows mitochondrial transport more than an N-terminal fragment [214]. Muhtt impedes axonal transport in its soluble form; aggregates can block axonal transport entirely producing axonal swelling and mitochondrial accumulation upstream of the block [214]. Beyond mitochondria, in vivo retrograde axonal transport of the exogenous marker molecule fluorogold along the striatonigral pathway was also impeded by the presence of mutant htt in YAC72 brain [132]. This interference with mitochondrial trafficking results in a paucity of ATP in synaptosomes harvested from Q150 mouse brains [1, 131]. Thus energy impairments in HD can result from a dearth of mitochondria at presynaptic release sites, limiting information flow and possibly at other sites of high energy demand without compromise of any enzymatic pathway.

**Mitochondrial dynamics**

Mitochondria in many HD model systems appear to be smaller than in normal tissue. In 6 mo old YAC128 striatum, in muhtt transfected cultured neurons and in fibroblasts from individuals with HD, mitochondria appear more fragmented than in controls [215, 216]. In YAC128 striatum compared to controls, a larger number of mitochondrial cristae occupy a smaller overall volume [216]. In these experiments, the mitochondrial fragmentation was attributed to an increased association between muhtt and DRP1, the protein responsible for fission, as dominant negative DRP1 prevented fragmentation [216]. Increased fragmentation was also observed in the immortalized STHdhQ111/Q111 cells who were also more sensitive to staurosporine-induced cell death than controls [120, 216]. Expression of dominant
negative DRP1 to prevent fragmentation or overexpression of OPA1 to prevent cristae remodeling, but not overexpression of MFN1 to promote fusion and elongation, rescued Q111 cells, lymphoblasts from HD donors and cultured YAC128 neurons from these phenotypes [216]. Fragmented or enlarged mitochondria were observed in about a quarter of cultured neurons from R6/2 striatum [110].

DRP1-dependent fission can become activated through multiple mechanisms [217, 218], several of which have been demonstrated to contribute to mitochondrial fragmentation in HD. In the STdh\textsuperscript{Q111/111} cells, fragmentation appeared to be triggered by increased calcineurin activation of DRP1 [216]. In a number of HD model systems, increased DRP1-induced fragmentation has been attributed to increased dopamine receptor driven calcineurin activity, increased S-nitrosylation of Cdk5 or direct association of muhtt with DRP1 [216]. Aberrant nitric oxide-induced S-nitrosylation of Drp1 also contributes to its activation and mitochondrial fragmentation in muhtt transfected primary neuronal cultures [219]. A selective inhibitor of DRP1 ameliorates mitochondrial fragmentation, restores mitochondrial membrane potential, and improves viability among Q111 cells and both HD patient fibroblasts and induced pluripotent stem cells differentiated into neurons. DRP1 inhibition also ameliorates symptoms in R6/2 mice [220]. Similar results were obtained from downregulation of p53, indicating that p53 might directly initiate DRP1-mediated fission in HD neurons [220].

Cristae disruption appears to be another consequence of DRP1 activation in HD [215, 216]. Cristae from neuronal mitochondria of 6 mo old YAC128 striatum had a higher density of cristae yet lower ratios for cristae to mitochondria surface area and volume [215]. Disorganized cristae were observed in Q111 cells and in primary cultures from neonatal YAC128 striatum [216]. However, fission and these cristae dilations are not incompatible with life as the Q111 cells proliferate and survive in vitro [221]. Further cristae disruptions triggered by apoptotic stimuli were accelerated in Q111 cells suggesting that cytochrome c release may become easier if spaces between cristae become enlarged [216]. In neurons, replenishment of peripheral energy generating capacity through fission of somal and axonal mitochondria is a critical homeostatic function [222]. These fissed mitochondria function normally [222, 223]. Ninety percent of anterogradely transported axonal mitochondria maintain a high membrane potential sensitive to ETC but not glycolytic antagonists [223]. In memory T cells, mitochondrial fusion and associated tight cristae promote OXPHOS while fission and cristae expansion promote glycolysis in effector T cells [224]. The effector T cells are viable, highly functional and metabolically active utilizing aerobic glycolysis and less efficient OXPHOS; such cristae changes are not severe enough to trigger immediate autophagy [224]. Striatal mitochondria express more of the DRP1 receptor FIS1 and are therefore more predisposed to become fissed [225]. If in addition, the observed similar cristae changes in HD neuronal mitochondria induce aerobic glycolysis [215, 216], this would be consistent with the proposed homeostatic shift in energy production towards glycolysis early in disease [65, 67].

Despite the increased incidence of smaller mitochondria in YAC128 striatum [215, 216], isolated YAC128 striatal mitochondria respire normally [108]. Thus these structural changes may increase neuronal sensitivity to apoptosis [216], but they do not do so by compromising ETC function. Other instances of normal mitochondrial bioenergetics function, despite altered shape, have been documented in neurons in which DRP1 activity has been downregulated. Selective depletion of DRP1 in DA neurons in vivo leads to swelling and loss of mitochondrial mass, but not membrane potential, so the degeneration under these conditions may be due to a lack of mitochondria rather than overt dysfunction of intermediate metabolism [226]. Similarly, knock out of DRP1 causes the normally elongated Purkinje cell mitochondria to become large and round with normal cristae morphology, normal membrane potential and ETC activity [227]. Even mitochondria from embryos of DRP1 null mice, which are embryonic lethal, maintain normal ATP levels when supplied with maternal energy sources, again suggesting that enzymatic activities are appropriate [228].

One contradiction remains to be untangled. AMPK, the master energy sensor, is also a regulator of mitochondrial fission. ETC enzyme inhibition of human osteosarcoma cells leads to increases in AMPK activity & consequent mitochondrial fission [229], a phenotype associated with increased uncoupling, a loss of nutrient storage and a decline in ATP production [3]. Since ETC inhibition appears to occur only late in HD progression, a similar mechanism is unlikely to account for mitochondrial fission early in disease progression. AMPK becomes activated by increased AMP/ATP ratios under conditions of cellular stress to increase energy production
and decrease energy-consuming processes [230]. For mild energetic stress including exercise or nutritional shortages, AMPK promotes mitochondrial fusion accompanied by increased oxidative phosphorylation [231]. For more severe stress triggered by radiation or toxic chemicals, AMPK promotes mitochondrial fission leading to autophagy and apoptosis [231, 232]. Exactly how AMPK signaling distinguishes between these divergent outcomes remains unclear. HD brains are characterized by normal energy generation at rest but an inability to produce sufficient energy in response to stimulation [66, 67]. Unlike the generic in vitro cell systems used to work out the above roles of AMPK, neuronal mitochondria are not elongated in HD, despite both the activation of AMPK [36] and the arguments that an inability to produce energy upon stimulation [66, 67] would be considered a mild stress, perhaps even milder than starvation. If AMPK activation is producing fission and autophagy, it must be doing so in a highly localized manner since death is not acute or rapid in HD. Also, AMPK may exert disease stage dependent effects [35].

To reconcile the metabolic results from experiments on mitochondrial dynamics with those on isolated mitochondria, one has to ask whether the isolated mitochondria represent fissioned or fused states. Newly isolated mitochondria are typically round and condensed when viewed by EM [108, 233, 234]. This results from the low ionic strength isolation media [233, 234]. When resuspended in osmotically isotonic potassium-containing medium, the condensed state rapidly changes to the orthodox configuration, with corresponding normally structured cristae. Addition of ADP to orthodox isolated mitochondria results in temperature-dependent condensation within 20 sec [234]. It is impossible to assert the original length or size of the mitochondria prior to isolation or if there are changes in size that occur as a result of isolation. The fission and fusion processes can occur within a minute while the centrifugations associated with isolation can take tens of minutes. Nevertheless, the normal ETC activity of isolated mitochondria from striatum and cortex of both the R6/2 and YAC128 mouse models of HD [108, 110] suggests that the original fused or fissioned state did not interfere with metabolic capacity of mitochondrial isolates.

Mitophagy

As with increased fission, loss of mitochondrial mass could of itself be sufficient to account for the energetic defects in HD. Indeed, mtDNA copy numbers were decreased in R6/2 striatum (but not cortex) and in STHdhQ111/111 cells [111, 120]. Mitochondrial mass is also decreased in HD induced pluripotent stem cells [5]. Mitochondrial clearance or mitophagy appears to be specialized forms of general autophagy, which is compromised in HD [235]. Muhtt may also act to alter mitochondrial clearance or mitophagy, although whether the effect is to hasten mitophagy or prevent it is currently disputed [5, 204, 236, 237]. In PC12 cells transfected with Q74 expanded htt and in STHdhQ111/111 cells, oxidized (and therefore inactivated) GAPDH associated with mitochondria targeting them to lysosomes but prevented internalization and mitochondrial degradation [204]. This process increased accumulation of damaged mitochondria and hence mitochondrial mass as evidenced by increased protein levels of VDAC, TOM20 and aconitase relative to enolase [204]. Overexpression of inactive GAPDH restored mitophagy, implicating muhtt as the species inhibiting this form of autophagy [204]. So caution should be applied when interpreting changes in mitochondrial enzyme expression if mitophagy is prevented. Valosin-containing protein (VCP) is a cellular ATPase associated with ER and mitochondrial degradation and autophagy. Muhtt on the mitochondrial outer membrane of STHdhQ111/111 cells and muhtt expressing induced-pluripotent stem cells binds to VCP, accelerating autophagosome formation or mitophagy [5]. This process is associated with loss of mitochondrial membrane potential, shortening of mitochondrial length in neurites, and increased cell death [5]. A synthetic peptide designed to inhibit muhtt-VCP interactions ameliorates these phenotypes plus motor phenotypes in R6/2 and YAC128 mice [5]. However it is not clear from either set of experiments whether the mitochondria are damaged prior to or as a consequence of GAPDH or VCP binding or recruitment. As an inducer of mitophagy, either protein could be causing the depolarization and fission. Thus it is possible that the energy generating pathways remain functional until mitophagy becomes initiated. Further clarification of the controversies surrounding mitophagy are needed to resolve its contribution to energetic defects in HD.

SUMMARY

This discussion has presented an updated, systemic view of shifts in brain energy generating
biochemical pathways in models of HD. The idea of a shift in energy status in HD is not new [133]. The proposed contextual homeostatic framework goes a long way towards reconciling the disparate findings across many laboratories. Regulatory feedback mechanisms control flux through the intertwined metabolic pathways, adjusting enzyme activities to meet local energy demands for normal function and the added burdens associated with accumulating mHtt. The ATP supply is sufficient for basal needs, but is limited in times of activation or stress. This loss of spare ATP slowly erodes protein clearance mechanisms, mitochondrial trafficking, and possibly even sodium clearance leading to accumulation of misfolded proteins and aggregates and perhaps hyperosmotic conditions [160]. These disruptions are not immediately lethal; rather they invoke regulatory compensations that shift over the course of disease adjusting to contextual factors.

Mitochondrial shape and size are intimately linked to whether glycolysis or oxidative phosphorylation is favored, adding another dimension to regulation of energy production. Elongated mitochondria favor oxidative phosphorylation and FAO while the fissioned state favors glycolysis and mHtt binds DRP1 promoting fission [153, 216, 224]. Movement of mitochondria within neurons and astrocytes may further limit energy supplies in distal processes. Early in disease, these homeostatic mechanisms may be sufficient to generate and provide the needed energy within the CNS. Transcriptional dysregulation [13, 14] would be expected to further compromise neuronal ability to generate energy, through down-regulation of CK expression or PDH activity or other metabolic enzymes. Slowly accumulating oxidative damage to individual metabolic enzymes would also strain capacities for flux through specific pathways, exacerbating energetic dysfunction as disease worsens. Glycolysis, TCA, ETC, and OXPHOS would be compromised by protein oxidation. Both TCA and ETC would be affected by loss of SDH subunit expression. Acetylation, (de)phosphorylation, or subsequent alterations in enzyme expression levels would further compromise fluxes. Oxidative or transcriptional limitations on transporters would limit energy substrates from entering cells or mitochondria, further restricting maximal energy production.

Compromise of the ETC appears not to be the principle or earliest metabolic change in HD pathogenesis. Overt loss of ETC function may not occur until late in HD progression, as seen in the post-mortem changes [238]. Early in disease flux through these energy generating pathways may shift towards glycolysis simply due to the rapidity of that response or due to the impediments in mitochondrial trafficking, propensity towards fission and increased mitophagy. Rather, compromise of glucose uptake facilitates glucose utilization to glycolysis and may possibly decrease flux through the PPP, limiting subsequent NADPH and GSH production needed for antioxidant protection. As a result, oxidative damage to key glycolytic and TCA cycle enzymes further restricts energy production so that while basal needs may be met, those of excessive stimulation may not be realized. Later more serious compromises may accumulate, e.g. decreases in SDH expression [238] and/or depression of glycolytic and TCA enzyme activities [65, 70, 72, 77]. Energy production may be further exacerbated by deficits in mitochondrial biogenesis or trafficking [2, 4]. The demonstration that in vivo upstream constrictions in metabolic flux may measurably restrict flux through ETC all the way to oxygen consumption [67] is a possibility not considered by many early energetic studies. Most bench metabolic studies are carried out under resting conditions, as opposed to those imposing a metabolic load, conditions that may better approximate in vivo conditions. The suggestion that substrate availability and transport into mitochondria may also be restricted needs to be explored further. These restrictions on energy production appear to be compensated for by stimulation of fatty acid oxidation and/or glutamate oxidation. The collective shifts in metabolic flux represent homeostatic compensatory mechanisms that maintain model organisms through presymptomatic stages and into symptomatic stages.

Only at end-stage disease would widespread deficits in ETC enzyme activity be expected, as observed in post-mortem regional human brain tissue. Some animal models may mimic this better than others. Surprisingly, in the most severe mouse model, R6/2, measurements of ETC activity, protein expression and ATP production diverge widely among laboratories, despite appropriate technical expertise [32, 33, 36, 66, 110, 111]. Other contextual factors may govern the rate of disease progression in these mice. The longevity of the R6/2 has been reported to vary widely between institutions and in response to environmental effects making it difficult to compare the degree of disease severity at a specific age [138, 239–242]. Stress in handling may be an example of one uncontrolled variable, especially when one considers that mice as a species can be
stressed by the presence of scents from male handlers [243]. Diurnal cycle disruptions and the time of day when experiments are performed may be other relevant factors [244, 245]. While difficult to demonstrate so many years after completion of these disparate studies, moving forward, stress-related factors should be carefully considered in the detailed execution of experimental design. The context of which muhtt fragments are active in distinct model systems influences the mechanisms of neuronal demise that ensues [11]. Similarly, contextual effects should influence how muhtt stresses mitochondrial functions.

**Future directions**

Appropriate questions to ask at this juncture are 1) can such homeostatic shifts be demonstrated all together in a single model system, and 2) what molecular or systemic events upend the homeostasis, sending the system into progressive decline. To answer the first question, a flux analysis approach should be applied longitudinally in several mouse models of HD. The second question addresses the relationship between metabolic dysfunction and clinical disease onset. Is the loss of homeostatic control attributable to mutant huntingtin itself or simply to normal aging? Intriguingly, an important energetic metabolite, NAD$^+$ has been proposed to be a key signal molecule for regulating the aging process [246]. NAD$^+$ is a key cofactor for sirtuins and for PARP reactions [246]. These important life-span regulating proteins compete for NAD$^+$, which acts as an ADP-ribose donor [246]. NAD$^+$ concentrations vary diurnally in mouse liver and decrease in muscle and liver over a lifespan [247, 248]. Hippocampal NAD$^+$ concentration drops after an ischemic insult and can be restored, along with behavioral recovery, by administration of nicotinamide mononucleotide, an activator of the NAD$^+$ salvage synthetic pathway [249]. In normal mouse aging, cortical, hippocampal and striatal NAD$^+$ (along with NADH and NADP$^+$) decrease between 18 and 24 mo as part of the age-associated overall increase in metabolite variability, termed metabolic drift [250]. NAD$^+$ has only been measured in ST$Hdh^{Q111/111}$ cells, where the NAD$^+$/NADH ratio increased in the polyglutamine expanded cells [94]. NADP$^+$ levels in dorsal R6/2 brain trended upward at 11 wk [33], but measures of NAD$^+$, NADH, NADP$^+$ and NADPH in other HD mouse brains have not been investigated. Connecting changes in longevity with metabolism may provide an important link for understanding the conflicting views of mitochondrial dysfunction in HD. Upregulating NAD$^+$ is recognized as being another energy boosting strategy without complete neuroprotective powers [246, 249]. However improving endogenous homeostatic responses in metabolic pathways may be a valid treatment strategy.

**Conclusion**

These results argue strongly that structural compromise of the ETC and the ability to generate ATP are not a direct cause for development of HD pathogenesis. The evidence that the metabolic rates through the various metabolic pathways change over the course of disease also argues that mitochondrial function and regulation are intact and operational over much of the lifespan of effected individuals and mice. The shifts in flux represent homeostatic responses to the changing transcriptional disruptions and increased energy demands for proteostasis. Temporal shifts in flux may occur in situ on a contextual basis without the need to invoke muhtt-induced changes in protein expression. Changes in mitochondrial dynamics and shape may be part of the homeostatic response as mitochondrial form follows function. Disruptions of mitochondrial biogenesis and trafficking further the need for homeostatic responses and add to the energy deficit. These progress until neurons cannot maintain sufficient energy production, eventually undergoing apoptosis. Energy production does however influence the time course of disease in that an increase in GLUT3 copy number delays onset of human disease [59]. Exogenous drugs that boost energy production or upregulate metabolic pathways ameliorate disease phenotypes or extend the lifespan of mouse models. Such interventions prolong the homeostatic compensation but do not address the underlying causes of accumulating muhtt. HD has been described as a disease in which adaptive transcriptional homeostatic mechanisms are disrupted by the polyglutamine expansion [14]. A similar homeostatic perspective should be applied to brain metabolism in HD.

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CONFLICT OF INTEREST

The author does not have any conflicts of interest.

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