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## The New Neurometabolic Cascade of Concussion

**Christopher C. Giza, M.D. and David A. Hovda, Ph.D.**

Division of Pediatric Neurology, Department of Pediatrics, Mattel Children's Hospital-UCLA, Los Angeles, CA

Department of Neurosurgery, David Geffen School of Medicine at UCLA, Los Angeles, CA

Interdepartmental Programs for Neuroscience and Biomedical Engineering, UCLA, Los Angeles, CA

Department of Medical and Molecular Pharmacology, UCLA, Los Angeles, CA

### Abstract

Since the original descriptions of post-concussive pathophysiology, there has been a significant increase in interest and ongoing research to study the biological underpinnings of concussion. The initial ionic flux and glutamate release result in significant energy demands and a period of metabolic crisis for the injured brain. These physiological perturbations can now be linked to clinical characteristics of concussion, including migrainous symptoms, vulnerability to repeat injury and cognitive impairment. Furthermore, advanced neuroimaging now allows a research window to monitor post-concussion pathophysiology in humans noninvasively. There is also increasing concern about the risk for chronic or even progressive neurobehavioral impairment after concussion/mild TBI. Critical studies are underway to better link the acute pathobiology of concussion with potential mechanisms of chronic cell death, dysfunction and neurodegeneration. This “new and improved” paper summarizes in a translational fashion and updates what is known about the acute neurometabolic changes after concussive brain injury. Furthermore, new connections are proposed between this neurobiology and early clinical symptoms as well as to cellular processes that may underlie long term impairment.

### Introduction

One of the hallmarks of concussion is that neurological signs and symptoms are imparted after biomechanical force to the brain in the absence of macroscopic neural damage<sup>1</sup>. In general, this has been interpreted as a result of predominantly functional or microstructural injury to neural tissue. Functional injury can refer to perturbations of cellular or physiological function including but not limited to ionic shifts, metabolic changes or impaired neurotransmission. Microstructural injury refers to physical changes not readily evident on CT scanning, but now detectable through advanced imaging (such as diffusion tensor imaging – DTI – for delineating signs of axonal injury). An important additional characteristic of sports-related concussion is the potential for repeated mild traumatic brain

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Corresponding Author: Christopher C. Giza, M.D., 300 Stein Plaza, 531 Wasserman, UCLA Department of Neurosurgery, Los Angeles, CA 90095, cgiza@mednet.ucla.edu.

injuries (TBI) over the course of an athletic event, a season or even a lifetime. The basic neurobiology of concussion/mild TBI has been elucidated in animal models, is increasingly corroborated in human studies and has been described as a neurometabolic cascade of events that involves bioenergetic challenges, cytoskeletal and axonal alterations, impairments in neurotransmission and vulnerability to delayed cell death and chronic dysfunction<sup>2,3</sup>. This review will use a translational approach to describe advances in our understanding of the underlying neurobiology of concussive injuries, with particular emphasis on linking the neurometabolic cascade to clinical characteristics as well as on new connections being made between acute post-concussion pathophysiology, long-term biological changes and chronic sequelae.

## Acute Pathophysiology (Figures 1 and 2)

### Ionic flux and glutamate release

Early studies demonstrated that biomechanical injury results in ionic flux and hyperacute indiscriminate glutamate release<sup>4,5</sup>. Potassium efflux, and sodium and calcium influx, occur due to mechanoporation of lipid membranes (creation of sublethal defects via traumatic insult) at the cellular level. Initial ionic flux and depolarization can then trigger voltage- or ligand-gated ion channels, creating a diffuse ‘spreading depression-like’ state that may be the biological substrate for very acute post-concussive impairments.

### Energy crisis

In an effort to restore ionic and cellular homeostasis, ATP-requiring membrane ionic pumps shift into overdrive, causing hyperglycolysis, relative depletion of intracellular energy reserves and increase in ADP<sup>6</sup>. In very early phases, this increased demand for energy occurs in a setting of normal or reduced cerebral blood flow, resulting in an uncoupling, or mismatch, between energy supply and demand.

Intracellular calcium flux, which occurs early and may persist longer than other ionic disturbances, is accommodated by sequestration of calcium into mitochondria. However, this short-term solution can then result in mitochondrial dysfunction, exacerbate problems with oxidative metabolism and worsen the cellular energy crisis.

In addition to the acute energy perturbations, intracellular redox state is altered. This puts additional stress on the system by generative damaging free radicals and shifting metabolic pathways that can trigger longer-lasting impairments and set the stage for vulnerability to repeated injury, which is particularly relevant for the clinical setting of sports-related concussion.

After an initial period of hyperglycolysis and metabolic uncoupling, glucose metabolic rates go into a state of impaired metabolism that can last up to 7-10 days in adult animals, and is associated with behavioral impairments in spatial learning<sup>6,7</sup>. The duration of this hypometabolic period appears to vary with age, with younger animals showing shorter periods (3 days) of impairment<sup>8</sup>. Post-traumatic changes in metabolism may be mediated by altered gene expression<sup>9</sup> and enzyme/transporter regulation<sup>10</sup>.

### **Cytoskeletal damage**

Biomechanical forces imparted onto neurons and glia can damage the delicate and complex microstructural components including dendritic arbors, axons and astrocytic processes. Following intra-axonal calcium flux, neurofilament side-arms can be phosphorylated and collapse, resulting in loss of structural integrity in axons<sup>11</sup>. Microtubule disruption due to axonal stretch can interfere with bi-directional axonal transport, potentially isolating the synapse, diminishing normal neurotransmission and in severe cases, result in axonal disconnection. Intra-axonal calcium flux results in proteolytic damage to subaxolemmal spectrin and other cytoskeletal components<sup>12</sup>. More recent studies have indicated that cytoskeletal anchor points in the cell membrane, mediated through proteins such as integrins, may represent a primary molecular target of traumatic injury in both neurons and vascular cells<sup>13,14</sup>.

### **Axonal dysfunction**

Axons prove to be particularly vulnerable to biomechanical stretch. Early studies showed increased axolemmal permeability after experimental TBI<sup>15</sup>. As mentioned above, damage to the neurofilaments and microtubules leads to axonal dysfunction and potential for disconnection. While axonal disconnection was once thought to invariably lead to demise of the neuronal cell body, *in vivo* animal studies have shown that perisomatic axotomy can occur with atrophy and shrinkage of the neuron, without cell death<sup>16</sup>. It is assumed that this type of severely damaged neuron is not capable of normal function. Some studies have suggested that dietary supplementation of particular fatty acids may mitigate, or at least alter the timeline of, axonal damage after experimental TBI<sup>6</sup>. *In vitro* stretch injury models demonstrate post-stretch axonal undulations and beading – in some cases these would recover, but in others the axonal damage was lasting<sup>17,18</sup>.

In addition, recent studies using fluid percussion injury centered over the white matter tracts of the corpus callosum indicate that unmyelinated axons are more vulnerable to injury (and show greater resultant impairment of electrophysiological function) than myelinated fibers<sup>19</sup>. This finding, although not replicated in a comparative study of immature animals, has developmental implications, as it is known that myelination of axonal tracts is one of the ongoing maturational changes that occur in the young brain.

Recent studies of repeat mild TBI in the immature brain have demonstrated white matter damage and associated cognitive impairments<sup>20</sup>. This is consistent with studies in adult animals that have also shown more prominent axonal injury with repeated mild TBI<sup>21</sup>.

### **Altered neurotransmission**

Changes in ligand-gated excitatory and inhibitory neurotransmission have been reported after *in vivo* experimental TBI. Alterations in glutamate (N-methyl-D-aspartate NMDA) receptor subunit composition and function have been reported following TBI in both the immature<sup>22</sup> and mature brain<sup>23-25</sup>. Functional consequences of NMDAR subunit change in developing rat pups have been described as interfering with normal developmental plasticity, electrophysiology and memory<sup>26</sup>. In adult rats, NMDAR changes have also been associated with functional alterations including differential patterns of calcium flux<sup>25</sup>,

immediate early gene activation and phosphorylation/activation of downstream signal transduction molecules such as CaMKII (calcium/calmodulin-dependent protein kinase II), ERK (extracellular signal-related kinase), CREB (cAMP response element binding protein) and BDNF (brain derived neurotrophic factor) <sup>23,27</sup>.

Post-injury excitatory-inhibitory balance can also be upset by changes in inhibitory neurotransmission involving GABA and its receptors. Studies have consistently shown dropout of hilar GABAergic interneurons following lateral fluid percussion injury (FPI) in adult rats <sup>28,29</sup>. In a recent study of post-TBI augmentation of fear-based learning, a putative model for post-TBI vulnerability to develop anxiety disorders and/or PTSD, decreased levels of GAD67, the biosynthetic enzyme for GABA, have been reported in the amygdala <sup>30</sup>. The amygdala is a critical structure implicated in the development of aberrant fear-based learning in animal models <sup>31</sup> but also in some human studies <sup>32</sup>.

### Inflammation

Somewhat overlooked in acute studies of mild TBI, changes in inflammatory markers have been well-reported in more severe TBI for some time <sup>33</sup>. Following TBI there is an activation and infiltration of microglia, particularly in cortical impact models but also after less severe models such as fluid percussion <sup>34</sup>.

More recent studies suggest that inflammatory changes are also triggered by mild TBI. In both adults and immature rats, microarray studies report extensive upregulation of cytokine and inflammatory genes after TBI <sup>9,35</sup>. Microglial activation after adult FPI, with or without exposure to the pesticide paraquat, has been associated with damage to the substantia nigra and implicated in the increased risk for parkinsonism after TBI <sup>36</sup>. Similar neuroinflammatory changes have also been reported subacutely in the absence of behavioral deficits in a different study <sup>37</sup>. A theory relating glutamate release and activation of immune receptors to oxidative stress and potentially later cellular injury has been proposed and termed “immunoexcitotoxicity” <sup>38</sup>.

### Cell death

Models of mild TBI have generally shown little cell death, even in settings where measurable cognitive impairments are described <sup>20,39,40</sup>. However, the consequences of repeated mild TBI are less well described – functional impairments appear to be greater than single TBI in most models <sup>21,41,42</sup>, raising the possibility that there may be longer term structural changes, although studies looking at chronic axonal injury and/or atrophy are lacking.

It is unclear whether chronic structural changes may evolve over longer time courses after even a single mild injury. However, following a single moderate/severe FPI in adult rats, ongoing cerebral and hippocampal atrophy have been described <sup>43</sup>, as have chronic losses in dopaminergic neurons in the substantia nigra <sup>36</sup>. Chronic evolution of injury has also been described after a single controlled cortical impact (CCI) injury in adult animals <sup>44,45</sup>. Immature (P7) mice show late appearance of cognitive deficits and tissue loss after a single CCI <sup>46</sup>, but other studies using a single FPI in developing rats (P19) showed no significant hippocampal or cortical neuronal loss <sup>39</sup>.

## Pathophysiology Meets Symptoms (Table 1)

While studies proving associations between the following pathophysiological changes after concussion/mild TBI remain to be conducted, it is tempting to speculate on how acute neurobiological change results in the clinical symptoms associated with concussion. Several reasonable connections can be made, and are presented in Table 1.

### Ionic flux and migraine

The widespread ionic flux described after FPI has often been described as ‘spreading depression-like’, and the original description of the spreading depression of Leao was in the context of migraine<sup>47</sup>. In fact, DC potential shifts have been described in both experimental TBI<sup>48,49</sup> and with direct electrocorticography after more severe human TBI<sup>50,51</sup>.

An additional intriguing connection that suggests a relationship between ionic compensatory mechanisms and cerebral responses to concussion is the description of significant mental status change and evidence for cerebral edema after mild TBI in the pedigree of a family with an ion channelopathy normally associated with familial hemiplegic migraine<sup>52</sup>. In this description, the CACNA1A mutation was associated with several clinical phenotypes, including an exaggerated response to mild TBI in some family members. There are an increasing number of ion channelopathies being identified, and it is interesting to speculate whether genetic factors such as these may underlie the ‘glass jaw’ phenomenon whereby some athletes are more readily symptomatic after mTBI, or whether this type of genetic vulnerability underlies the malignant cerebral edema seen in cases of ‘second impact syndrome’.

Finally, there is considerable overlap of typical post-concussion symptoms with those commonly described in migraine, particularly headache, nausea, vomiting, photophobia, phonophobia and dizziness. In fact, some studies indicate the reporting of headache symptoms or a prior history of migraine as potential risk factors for prolonged recovery or more severe symptoms after concussion<sup>53</sup>.

### Energy crisis and vulnerability

Clinically it is known that a history of prior concussion/mTBI is a strong risk factor for subsequent concussion, and that in prospective studies, this risk appears greatest in the first 10 days post-injury<sup>54-56</sup>. There are many factors that may play into this increased clinical risk, but biological vulnerability due to ongoing energy crisis is one that can be addressed by the current practice of delaying return to contact risk. The challenge here is identifying the duration of this period of metabolic vulnerability, as it likely varies between concussed individuals.

Human studies using magnetic resonance spectroscopy (MRS) have shown reductions in the metabolite N-acetylaspartate (NAA) in frontal white matter of concussed adults compared to normal controls<sup>57,58</sup>. Over 30 days, NAA levels returned to normal, except in individuals who sustained a second concussion, in whom NAA did not recover fully until 45 days after the initial injury<sup>58</sup>. Such findings support the concept of metabolic vulnerability after concussion. However, in these studies, the depressed NAA levels did not correlate well with

reported symptoms, and thus the clinical significance of post-concussion NAA remains uncertain.

Animal studies have shed some light on this. In a weight drop model in adult rats, the greatest perturbations of neurochemical markers of metabolic stress occurred when injuries were separated by 3 days<sup>10,59</sup>. Repeated concussive injuries in adult mice showed worsening of neurocognitive function and traumatic axonal injury when injuries were spaced out by 3 or 5 days, but not when injuries were separated by 7 days<sup>21</sup>.

In a closed head impact model in juvenile rats, the period of reduced glucose metabolism post-injury was associated with impairments in working memory and generally lasted about 3 days. When a second injury occurred within this period of impaired glucose metabolism (post-injury day 1), the severity of both hypometabolism and memory impairment was greater. If the second injury occurred after full metabolic recovery from the first injury (post-injury day 5), the two injuries acted like single, separate injuries<sup>42</sup>.

Overall, these investigations indicate that the timing of repeated injuries may have significant consequences on the severity of post-concussion pathophysiology and subsequent cognitive disturbances. They also indirectly support the current clinical management protocols of removing concussed athletes from contact risk immediately to allow time for biological recovery. Further work is needed to identify what biomarker can be used to determine the time of recovery for individual patients – symptoms, cognition, balance, reaction time and neuroimaging have all been proposed. It is likely that the duration of time for recovery in humans will be longer than that described in the animal models.

### **Axonal dysfunction and slowed cognition**

White matter injury and damage to neural networks has been associated with cognitive impairments after moderate and severe TBI. Diffuse axonal injury (DAI) is traditionally described as a pathological finding of damage to white matter tracts associated with frank axonal disconnection and microhemorrhages, often seen post-mortem after severe TBI<sup>60</sup>. Furthermore, it is recognized that after TBI, axonal injury also occurs on a microscale, with mechanoporation, stretching and beading of individual axons, disruption of axonal transport and axonal blebbing, as well as disconnection. The complete spectrum of axonal pathophysiology is sometimes termed traumatic axonal injury (TAI), to distinguish it from the more classical pathological description of DAI<sup>61</sup>.

For mild TBI and concussion in humans, there is little pathological data<sup>62</sup>, but advanced MRI using diffusion tensor imaging (DTI) has been used to measure changes in directionality of water diffusion that may be indicative of white matter disruption. Fractional anisotropy (FA) one such measure, with higher FA values indicating more water diffusion in a particular direction and lower FA values indicating less directionality. Increased FA values have been reported in corpus callosum early after mild TBI (6 days) in adolescents, and these values were correlated with a validated symptom scale<sup>63</sup>. Early increases in FA were also reported in a cohort of adolescents after concussion but no correlation was found between the DTI changes and cognitive function<sup>64</sup>. Recent studies have reported changes in

DTI measures and cognitive testing after a period contact/collision sports participation in individuals with no diagnosed concussion<sup>65,66</sup>.

For more severe TBI, damage to the white matter strongly correlates to burden of neurocognitive impairment. FA generally is decreased in major fiber tracts (corpus callosum being most studied), and white matter disruption is associated with burden of neuropsychological deficits using standard testing<sup>67,68</sup>.

Overall, data from DTI studies of mild TBI has been more variable, with some acute studies showing increased FA<sup>64,69-71</sup> and some showing decreased FA<sup>65,72,73</sup>. It is likely that multiple parameters can affect the results of DTI studies, including timing after injury, age of subjects, severity of injury, region of brain analyzed as well as the specifics of the methods used<sup>74</sup>. Thus, while DTI appears to be a very sensitive imaging modality to study concussion/mTBI noninvasively, its use for routine clinical management of individual patients is still premature.

Again, animal studies provide some additional perspective. Using a mouse model of more severe TBI - controlled cortical impact - correlations between DTI measures and histopathology were elegantly described across a time range from hours up to one month post TBI<sup>75</sup>. These studies show that DTI is more sensitive than conventional MRI at detecting white matter injury after TBI. They also shed some light into the changing nature of DTI signal as the TBI evolves over time. Early changes (<24 hrs) included maximal staining of axonal varicosities with amyloid precursor protein (APP) and neurofilament light chain markers, while later time points (4 days to 1 month) showed maximal staining for gliosis (GFAP - glial fibrillary acidic protein).

Rodent models of mild repeated TBI have shown impairments in spatial learning and memory in the absence of overt histological damage or cell death<sup>41</sup>. Increased APP was most prominent in animals with repeat closed head injury separated by 3 days, and these animals also showed the greatest impairment in motor function and spatial learning, when compared to single injury animals or those with repeat TBI separated by 1 week<sup>21</sup>. Graded deficits in working memory after mild and repeat closed head injury (24 hours apart) have been reported along with increases in APP staining in corpus callosum and subcortical white matter<sup>20</sup>.

Taken together, these studies show that cognitive deficits occur after mild TBI in the absence of overt cell death. Graded degrees of axonal injury/dysfunction occur after mTBI and repeat mTBI, and can be associated with graded levels of neurobehavioral impairment. However, in mild TBI, the linkage between imaging measures of axonal damage and clinically measurable cognitive problems is still in evolution.

### **Altered neurotransmission, slowed cognition and reaction time**

Neurotransmitter and synaptic alterations are other mechanisms of neural dysfunction that occur in the absence of overt cell death and may contribute to acute and transient neurobehavioral impairments after concussion. Clinical studies show early post-concussion problems in attention and memory that generally recover over time; these have been

measured using simple sideline questions <sup>76</sup>, the Standardized Assessment of Concussion (SAC) <sup>77</sup>, paper and pencil neuropsychological testing <sup>78</sup> and computerized cognitive testing <sup>79</sup>. Slowed reaction time has been measured in conjunction with computerized tests as well as with a simple ‘stick-drop’ device <sup>80</sup>.

Determining the physiological correlate of impaired post-concussion cognition has been attempted through the use of functional MRI (fMRI). Functional MRI measures changes in blood oxygen level dependent (BOLD) signal driven by increases in neural activity associated with specific cognitive tasks. While there remains debate about the physiology of the BOLD signal <sup>81</sup>, evidence from animal models shows that up to 50% of the BOLD signal may be driven by excitatory glutamatergic neurotransmission <sup>82</sup>.

Human studies of fMRI activation following concussion have often used cognitive measures of working memory. Abnormal patterns of increased BOLD activation have been reported days and weeks after concussion/mTBI in adults <sup>83</sup> and older teens (average age 16.6y) <sup>84</sup>. These changes may serve as a biomarker of ongoing neural dysfunction or in one study, as a marker for prolonged recovery <sup>84</sup>. A more recent study of slightly younger teens (average age 14.5y) after concussion showed worse working memory associated with significantly reduced task-related BOLD activity <sup>85</sup>.

As mentioned above, animal studies have demonstrated TBI-induced deficits in excitatory neurotransmission, often involving the NMDAR <sup>22-25</sup>. Interestingly, in the immature brain, these impairments are associated with a loss of experience-dependent plasticity that results in lasting learning and memory deficits <sup>86,87</sup>. Even if a period of ‘recovery’ is allowed before stimulating experience-dependent plasticity, restoration of memory functions is only partial <sup>88</sup>. Ongoing studies are aimed at developing mechanism-based interventions to help restore these impairments.

It is difficult to entirely tease out the contributions of axonal injury vs impaired neurotransmission to post-concussion cognitive and behavioral deficits. Quite likely both are involved, and determining the predominant mechanism in a given individual may depend on continued study into biomarkers to help identify those at risk for persistent impairments, and to determine interventions most likely to be effective.

Interestingly, as mentioned above, while post-traumatic metabolic changes in the immature brain appear to be shorter lasting than in adults <sup>8</sup>, axonal vulnerability <sup>19,20</sup> and impairments in plasticity <sup>86,87</sup> may be more prominent in the young brain. Additional work is necessary to determine on balance whether the young brain is more or less vulnerable to mild TBI, although some clinical evidence suggests, at least for concussion, recovery takes longer in high school athletes compared to collegiate or adult professional athletes.

## How much is too much? Acute-to-Chronic Pathobiology (Figure 2)

Evidence for the chronic sequelae of repeated mild TBI/concussion has long been associated with pathological studies of boxers <sup>89,90</sup>. More recently, case studies of contact sport athletes have shown both gross and microscopic pathology that has been attributed to repeated exposure to concussive blows <sup>91,92</sup>. In these more recent cases, deposition of



aggregated tau protein has been reported as the pathological hallmark of the condition now termed chronic traumatic encephalopathy (CTE). However, the waters surrounding our current understanding of CTE have been muddied by selection bias inherent to any case report study, an uncertain dose relationship between mTBI and pathology, concomitant associations between repeat concussion and other neurodegenerative conditions and the lack of a unifying biological mechanism to translate the acute (but generally recoverable) pathophysiology of concussion into a chronically progressive disorder.

### **Energy crisis, protease activation and cell death**

Acute impairments of cellular metabolism have described problems in both glycolytic and oxidative pathways (minutes to days, see above). Excess calcium flux is sequestered in mitochondria but then results in diminution of oxidative capacity that can spill over into the subacute period (days-weeks). Contact/collision sports and military duty, among other professions, are distinct from most accidental mTBI because of the inherent goal of returning to contact risk activity. The occurrence of repeated mTBI prior to full recovery from a first mTBI may set in play more lasting metabolic perturbations that may then trigger activation of intracellular proteases and the cascade that leads to apoptotic cell death.

Human studies have not definitively demonstrated this progression, which can only be done using longitudinal study designs. However, it is well known that gross pathology seen in the brains of boxers and others exposed to repetitive concussions includes cortical and hippocampal atrophy, ventriculomegaly, cavum septum pellucidum and other findings strongly suggestive of chronic cell death<sup>93</sup>. More recent studies are beginning to use advances in quantitative MRI and longitudinal study designs to examine progressive atrophy within subject after mild TBI<sup>94</sup>.

Animal studies support the concept that more severe TBI can result in chronic evolution of injury that results in ongoing cell death and atrophy out to 1 year<sup>45,95</sup>. Even in a pediatric model of moderate-to-severe experimental TBI, progressive cell death and volume loss, associated with emergence of additional cognitive decline, has been reported over 1-3 months post-TBI<sup>46</sup>. Longitudinal studies of brain atrophy after experimental repeat mild TBI/concussive injury remain to be done.

### **Altered protein degradation and toxic accumulation**

In addition to the effects of chronic energy impairment as a trigger to protease activation and apoptotic cell death, it is well known that normal cellular protein homeostasis depends upon a functioning system of protein degradation. This system requires energy, ubiquitin and a protein complex called the proteasome<sup>96</sup>. There are many examples in neurodegenerative disease of cellular oxidative stress leading to oxidatively damaged proteins that can affect metabolic enzymes and/or the ubiquitin-proteasome system. This could then result in the accumulation of abnormal/toxic proteins<sup>97</sup>. These mechanisms have been implicated in Alzheimer disease, tauopathies, synucleinopathies (Parkinson disease) and ALS, to name a few, and it is not surprising that these links are now being made in TBI. Early and prolonged proteasomal activation has been proposed based upon immunohistochemical findings after severe human TBI<sup>98</sup>.

Some experimental studies have identified molecular and functional changes in cortical and hippocampal proteasomes after TBI<sup>99</sup>. This suggests proteasome dysfunction is a viable mechanism as a precursor to post-traumatic neurodegeneration, although these studies did not specifically measure for accumulation of toxic or abnormal proteins. A more recent study linked abnormalities in proteasomal molecules with enhanced apoptosis after TBI<sup>100</sup>.

Accumulation of abnormal proteins has also been reported in animal models of TBI under multiple circumstances. Wild-type rodents show accumulation of oligomeric and phosphorylated tau after parasagittal fluid percussion injury<sup>101</sup> and blast neurotrauma<sup>102</sup>. Interestingly, deposition of amyloid or tau was *not* seen at chronic time points in a study of repetitive mild TBI, even though the animals showed cognitive impairments and astrogliosis<sup>103</sup>. Transgenic animals (3xTG-ApoE4) have also been reported to show increased accumulation of abnormal proteins after TBI<sup>104,105</sup>.

Aggregation and progression of toxic molecules has recently gained traction in a series of articles suggesting that damaged tau oligomers, not neurofibrillary tangles, are the triggering molecules for spread of tau pathology in Alzheimer disease<sup>106,107</sup> (for review see<sup>108</sup>). Elevated levels of extracellular tau have been reported after severe human TBI, and may hypothetically initiate the process of progressive tauopathy<sup>109</sup>.

Based upon these data, the mechanism of impaired protein homeostasis and degradation after TBI is plausible. Confirmatory studies remain to be conducted. Such a mechanism may likely involve oxidative damage to metabolic or proteasome molecules. There is evidence that proteins such as amyloid and tau, which accumulate in known neurodegenerative disorders, also show perturbations after both clinical and experimental TBI.

### Chronic axonal dysfunction

As discussed above, axonal dysfunction can result in impaired cognitive functioning and slowed reaction times, either through slowed conduction, damage to cerebral networks or deficits in neurotransmission. These mechanisms may be detected early after mTBI using DTI and fMRI<sup>64,83</sup>.

DTI has also been used to detect white matter abnormalities subacutely and chronically after mTBI. One recent study used DTI to examine military personnel evacuated from Afghanistan and Iraq for TBI, and found persistent changes in multiple regions<sup>110</sup>. Even in a civilian youth cohort with concussions, changes in DTI were detected early after injury while the subjects were symptomatic, but then showed similar changes at 4 months post-injury, a time when most were asymptomatic. The clinical significance of these DTI signal changes in a subacute/chronic setting is only beginning to be understood, but these white matter alterations have been associated with post-traumatic stress disorder<sup>111</sup> and depression<sup>112</sup> in different cohorts. Perhaps equally concerning are recent reports of abnormal DTI measures in athletes subjected to repeated mild head blows (heading in soccer) even without a diagnosis of concussion<sup>65</sup>.

While axonal disconnection after severe TBI can certainly result in cognitive impairment, it is uncertain how much post-concussion white matter pathology is due to disconnection vs

alterations in axonal physiology<sup>18</sup>. Furthermore, the idea that axotomy results in the inevitable death of the neuron has been challenged by evidence showing some neurons survive axonal disconnection but may be rendered dysfunctional<sup>16</sup>. What also remains uncertain is the ability of the damaged axon to recover, or the time frame of events that dictates recovery vs disconnection. Recent evidence has focused on the possibility for persistent axonal degeneration following TBI, perhaps even a single TBI<sup>17,113</sup>. As mentioned earlier, myelination also appears to protect axons from trauma, with unmyelinated axons more vulnerable<sup>19</sup>. These mechanisms of chronic axonal damage may then be particularly problematic in settings of repeat mTBI without adequate time for recovery between injuries, immature myelination or individuals with genetic vulnerabilities.

## Summary

Our understanding of the pathophysiology of concussion has expanded significantly in the last decade. Many of the acute physiological changes originally described have received additional support in newer models of mild repetitive TBI and with the use of advanced noninvasive neuroimaging. The contributions of oxidative stress, impaired axonal transport and altered neurotransmission are now clearly linked to the initial ionic fluxes, indiscriminate glutamate release and metabolic uncoupling. Furthermore, it is increasingly possible to see links between the pathophysiology of concussion and the early clinical signs and symptoms; there is great potential that, as the time course of these cellular events is better elucidated, this information will translate to a clearer idea of the timing of increased risk for repeated concussion in patients. Other accelerated research efforts into the ‘new’ neurometabolic cascade of concussion include studies examining causal linkages between the acute pathobiology and chronic progressive changes that may result in lasting impairments. While still early, untangling the connections between acute and chronic pathophysiology of concussion holds the promise for better prevention of repeated injury and mechanism-based therapies to interrupt the progression to persistent deficits or neurodegeneration.

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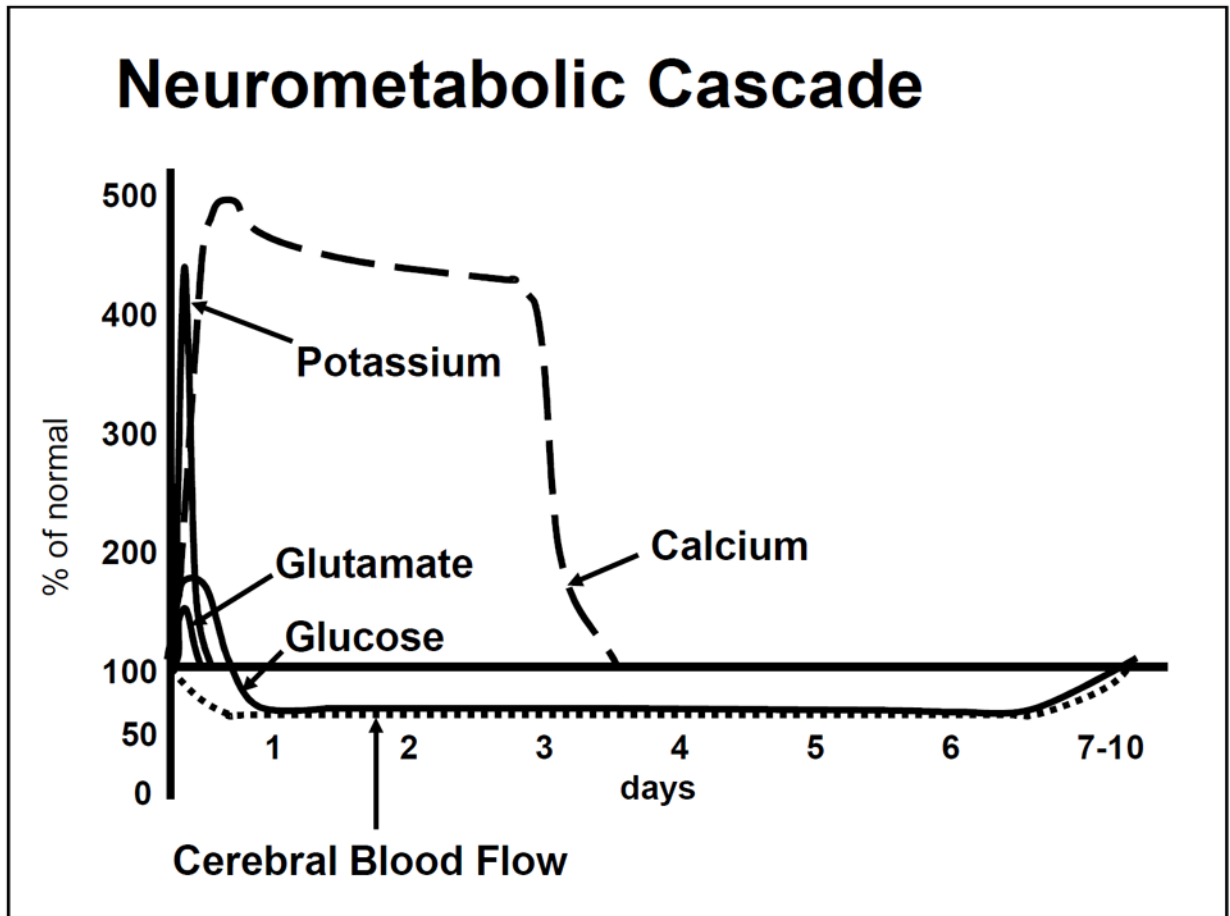
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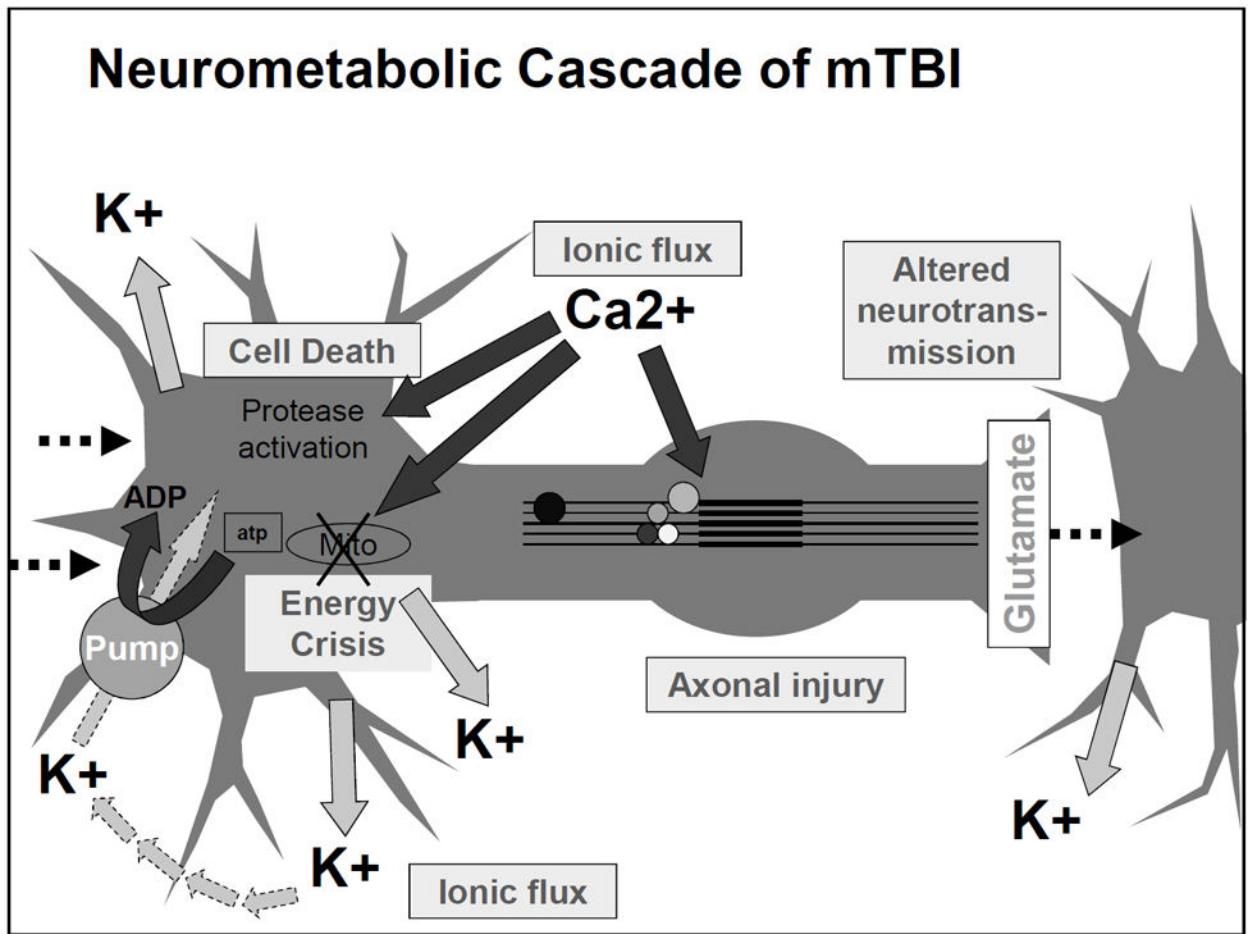
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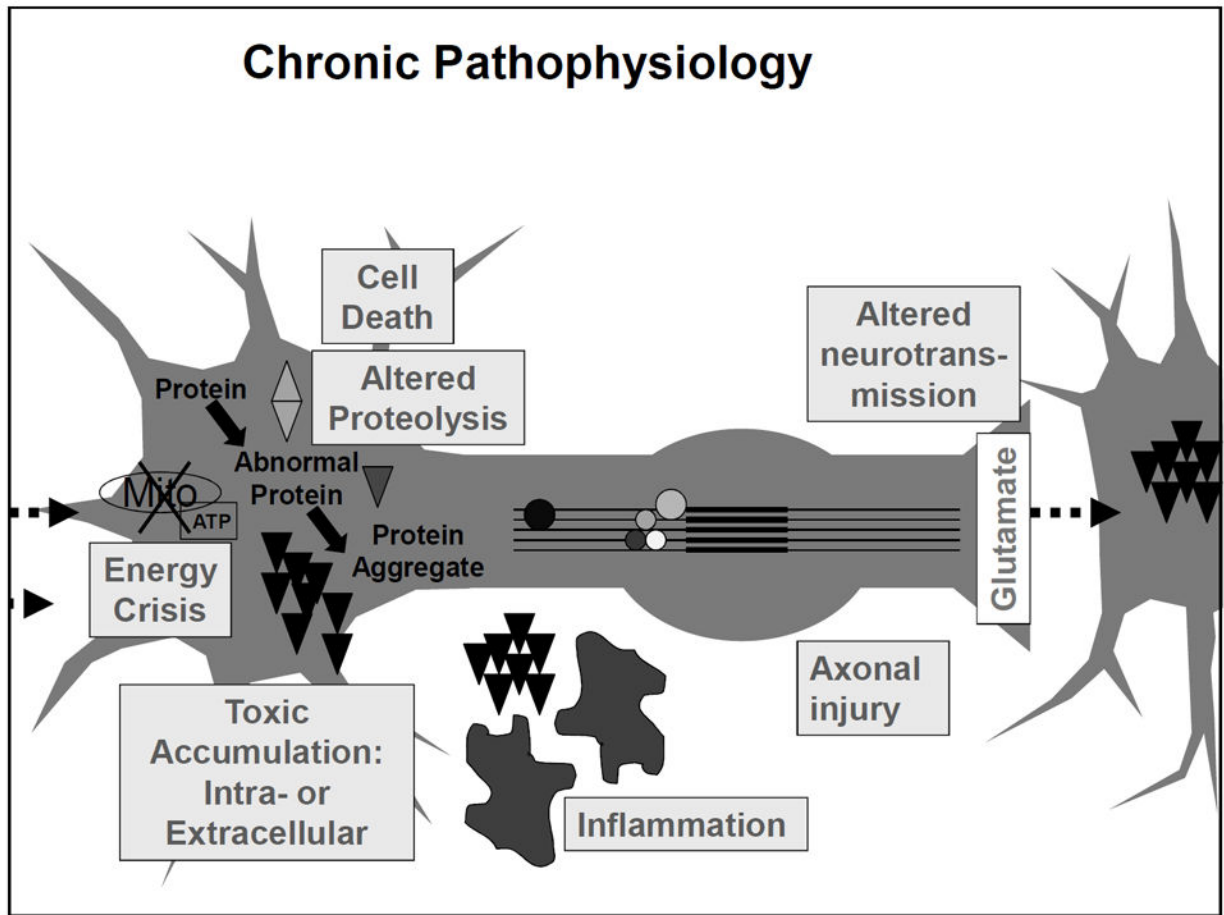
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**Figure 1.**  
Time course of the neurometabolic cascade of concussion.



**Figure 2.**  
Diagram of the acute cellular biological processes occurring after concussion/mild TBI.



**Figure 3.** Diagram of linkages between acute post-concussion pathophysiology and mechanisms for chronic impairment and possibly neurodegeneration.

**Table 1**

Physiological perturbations after concussion and proposed clinical correlates.

<b>Post-TBI pathophysiology</b>	<b>Acute symptom / clinical correlate</b>
Ionic flux	Migraine headache, photophobia, phonophobia
Energy crisis	Vulnerability to second injury
Axonal injury	Impaired cognition, slowed processing, slowed reaction time
Impaired neurotransmission	Impaired cognition, slowed processing, slowed reaction time
Protease activation, altered cytoskeletal proteins, cell death	Chronic atrophy, development of persistent impairments

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