The prevailing attitude in sports culture was once rather dismissive of concussions, viewing them as minor injuries without consequence. This mindset has undergone a major overhaul, particularly in the last 10–20 years, and concussions are now taken much more seriously, in part because of the forced retirement of several high-profile athletes. The effects of a single concussion are rarely long-term owing to the spontaneous resolution of symptoms and better management strategies; however, long-term changes ranging from subtle electrophysiological alterations in cognitive function to persistent or more slowly resolving symptomatology as well as neurocognitive changes on reliable neuropsychological tests are detectible, particularly after multiple concussions. Furthermore, anatomical changes have also been documented. In this review, across a variety of subclinical and clinical findings pertaining to the spheres of cognition and motor function, we discuss the relevant differences observed in the chronic postinjury phase in active as well as retired athletes who have sustained multiple concussive blows.

Cognitive Function

Cognitive Function in Young, Active Athletes With Multiple Concussions

Neuropsychological assessment, the gold standard in concussion diagnosis and management, outside the acute postinjury phase has revealed mixed results with respect to the long-term consequences of concussion in athletes. Iverson and colleagues found that high school athletes with a history of 1 or 2 prior concussions did not demonstrate any neuropsychological deficits 6 months after injury relative to athletes who had not suffered any concussions. In contrast, a study of amateur soccer players showed that those with a history of concussion performed more poorly on neurocognitive tests. In another study, Collins and colleagues reported lowered baseline testing in college athletes with a history of concussion, although this was in conjunction with a learning disability, which may imply a detrimental synergistic relationship between concussions and impaired learning abilities. Another study revealed that professional soccer players, relative to elite noncontact sports athletes, demonstrated residual neuro-
which may be elevated in the chronic postinjury phase. To normalize by 6 months postinjury, and myoinositol, include diminished levels of both glutamate, which seem nonconcussed athletes. Other neurometabolic alterations consistently diminished in concussed athletes relative to the integrity of white matter by using the diffusive proper-

ing is a noninvasive means of investigating the structural changes. The diffusion tensor imaging literature to date suggests the number of previous concussions is the prevail-

ing finding in young athletes with multiple concussions tested several months after their last injury. Authors of a more recent study reported persistent neurophysiological measures of altered short-term memory retention in athletes with more than 3 concussions, as revealed by a significantly suppressed sustained posterior contralat-

eral negativity waveform elicited by a visual short-term memory paradigm. Altered neurophysiological me-

sures of performance monitoring was also found in athletes with multiple concussions, suggesting that recurrent concussions impair error processing mechanisms during cognitively engaging tasks. Authors of the expert panel from the last symposium on return-to-play protocols is the inability to relate these persistent ERP components alterations to either self-reported cognitive symptoms or behavioral performance alterations of the corpus callosum, posterior limb of the internal capsule, superior longitudinal fasciculus, and superior frontooccipital fasciculus to name a few. These alterations oc-

cur in part due to the highly variable nature of how concussions are sustained and in part due to individual fac-

tors. Diffusion tensor imaging studies have yet to focus on differences between athletes with single and multiple injuries, although Henry et al. found that the number of concussions did correlate with the number of brain regions showing diffusion alterations. A more recent tech-

nique, such as susceptibility-weighted imaging, has yet to be used enough for its efficacy to be assessed, although early evidence suggests that it may have utility in a pedi-

atric population. It should be stressed that in all of the aforementioned techniques, the functional consequences for the metabolic or structural changes are unknown.

Subclinical Alterations in Cognitive Function

Event-related potentials are averaged electrical brain responses that allow one to determine the time course of higher level processes such as attention and memory up-

dating in the human brain. The ERP measure represents the averaged electroencephalography signal time-locked to a presented stimulus and consists of different compo-

nents labeled by their polarity (for example, P for positive and N for negative) and temporal range in milliseconds or by the ordinal number of major components. To date, most of the literature on ERPs and mild TBI has focused on the modulation of the classic P3 response as a result of concussion. The appeal of measuring the P3 component size and latency lies mostly in the ability to index changes in cognitive efficiency following a concussion. Evidence of enduring P3 changes in athletes who have suffered multiple concussions has abounded over the last decade. A P3 amplitude reduction exacerbated as a function of the number of previous concussions is the prevailing finding in young athletes with multiple concussions tested several months after their last injury. Authors of a recent study reported persistent neurophysiological evidence of altered short-term memory retention in athletes with more than 3 concussions, as revealed by a significantly suppressed sustained posterior contralat-

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on validated neuropsychological measures of memory and attention functions beyond the acute postconcussion phase. Another major limitation is the relatively poor specificity of these ERP components, although they are highly sensitive to the effects of sports concussions. For instance, alterations in P3 components have been found in various clinical populations including those with schizophrenia, alcoholism, depression, attention deficit disorders, epilepsy, Alzheimer disease, and others. The poor ERP component specificity is particularly problematic for concussed athletes whose P3 components alterations are not tied to observable or reported cognitive difficulties, as one might argue that factors extraneous to the damaging impact of sports concussions on brain functions, such as concomitant depressive symptoms, attentional difficulties, or sleep disturbances, could just as well have influenced the P3 size and latency. Moreover, without supporting evidence from prospective studies focused on changes in ERP components relative to concussed athletes’ own baseline measurements, we cannot determine whether these ERP components alterations are a consequence of sports concussions rather than a premorbid characteristic of athletes who are more at risk for sustaining sports concussions. Nevertheless, the consistent nature of ERP results across studies suggests that there may be subclinical alterations in athletes who have suffered multiple concussions.

Cognitive Function in Late-Adulthood Athletes With Multiple Concussions

While not typically disturbing daily life functions in young athletes, these subclinical alterations were linked to overt late-life clinical manifestations in a recent study conducted in late-adulthood athletes presenting with a history of remote concussions. Although not fully addressing the issue of cause and effect, evidence for the clinical significance of chronic P3 components alterations after sports concussions was revealed in a group of former athletes with concussions who had demonstrated reduced performance scores on neuropsychological tests that significantly correlated with P3a/P3b components amplitude attenuations. This finding confirms that concussion-related P3 components attenuations are associated with a worse performance on neuropsychological tests of episodic memory and executive function as athletes age and that the more the P3 components are suppressed, the greater the expected cognitive decline. The concept of “cognitive reserve” is very useful to help explain the resurgence of neuropsychological test performance decrements when athletes are tested more than 30 years postinjury. The central tenet of the cognitive reserve concept is that resilience to brain damage reflects how the brain uses its available resources to compensate for damage by optimizing performance through differential recruitment of brain networks and/or alternative cognitive strategies. Another important aspect of cognitive reserve is that damage from various environmental, developmental, and genotypic sources accumulates throughout a lifespan to reduce cognitive reserve. The progressive reduction in cognitive reserve via the accumulation of brain-damaging experiences and slowly waning health conditions throughout a lifespan characterizes the aging process and the accompanying decline in cognitive function. When applied to concussion findings, evidence suggests that young concussed athletes who showed persistent P3 components alterations can rely on their cognitive reserve, thus allowing the recovery of baseline performance levels on neuropsychological tests after the acute postconcussion phase had subsided. Some 1H-MRS evidence indicates continued metabolic alteration in asymptomatic athletes with concussed brain, while other data have shown that there is metabolic recovery. Although there is much work to be done to clarify if and how cognitive reserve plays a role in the compensation for and recovery from concussion, what seems more certain is that regardless of an asymptomatic state and a normal neuropsychological performance, subclinical differences persist in athletes who have sustained a concussion. Further, these differences are moderated by the number of concussions an athlete has sustained. The picture becomes much clearer when older former athletes are considered. When the chronic deleterious effects of sports-related concussions on cognitive function were combined with known age-related cognitive decline, the ensuing cognitive reserve of formerly concussed athletes, relative to that of nonconcussed athletes, could no longer maintain optimal performance levels on neuropsychological tests of episodic memory and executive functions selected for their sensitivity to early symptoms of MCI. Note that MCI is known as a transitional stage between normal aging and dementia, including Alzheimer disease. Of great clinical relevance, former athletes with concussions performed significantly worse than their nonconcussed counterparts on cognitive functions that have recently been shown to be affected in the very early stages of MCI. Most notably, episodic memory decline in former athletes with concussion relative to controls was found in the recognition component of the Rey-Osterrieth Complex Figure Test, and scores on the immediate and delayed recall components also tended to be lower. Interestingly, visual recognition memory impairments were recently found early in the course of MCI. The acute sensitivity of visual recognition memory tests to MCI was related to the distribution of neurofibrillary tangles over hippocampal structures, which is known as a core neuropathological hallmark of Alzheimer disease. The recent demonstration of strong correlations between visual recognition memory decline and 1H-MRS/volumetric MRI evidence of compromised hippocampal formation integrity in former athletes in late adulthood with remote concussions highlights important similarities between the pathophysiology of sports concussions and that of MCI. However, the presence of widespread cortical thinning together with elevated choline levels in dorsolateral prefrontal cortices not typically found in patients with MCI suggests that sports concussion in aging athletes exerts its own pathophysiological signature.

Motor Function

Motor Function in Young, Active Athletes With Multiple Concussions

While the recovery of cognitive impairments after sports concussions has drawn most of the attention from

Neurosurg Focus / Volume 33 / December 2012
the scientific community over the last few decades, the investigation of motor system abnormalities has recently come to the forefront of the sports concussion literature. Indeed, assessment of postural stability with various stances and surfaces was recently integrated into clinical practice to assist clinicians in determining when concussed athletes who experienced balance problems after injury could safely return to play. While postural stability has typically been found to return to baseline levels within a few days postconcussion on conventional measures of COP displacement, the recent addition of approximate entropy calculation as a nonlinear dynamic measure of postural control has revealed increased sensitivity to subtle physiological alterations associated with sports concussions. Indeed, athletes who resumed competition more than 9 months prior to testing still exhibited enhanced COP oscillation regularity according to an approximate entropy measure of postural control despite normal postural stability scores on conventional linear measures. Although the functional significance of enhanced COP oscillation regularity with regard to postural stability is still largely unknown, previous studies have suggested that it may represent an adaptive compensatory mechanism put forth by concussed athletes to achieve postural stability. Another possibility is that concus- sive injuries result in stiffened lower extremity musculature potentially related to excessive motor system excitability after sports concussions.

The application of TMS for sports concussion emerged mostly from postural imbalance data possibly involving motor system excitability abnormalities coupled with the TMS demonstration of altered cortical excitability of the M1 in the acute postconcussion phase. This relatively novel technique has been reliably used to detect preclinical covert indices of motor disorders. Among primary TMS measures, the CSP revealed unprecedented sensitivity to the subtle alterations in central inhibitory/excitatory mechanisms of the motor system in concussed athletes. The later part of the CSP has been attributed to activity of GABA<sub>B</sub>-mediated intracortical inhibitory sys- tems of the M1, whereas spinal inhibition contributes to its early part. Increased intracortical inhibition of M1, as reflected by prolonged CSP duration, was found in concussed athletes who had been asymptomatic on average 2 years prior to testing. In an earlier study, we prospectively examined the effects of recurrent concussions in a small group of athletes with multiple concussions who were retested using the same experimental protocol and found that the CSP was further prolonged after another concussion was sustained, suggesting that M1 intracortical inhibitory receptors may be particularly vulnerable to the effects of subsequent concussions. Based on these findings, it would appear that concussions alter the efficacy of GABA<sub>B</sub> receptor systems, perhaps contributing to rendering the brain more vulnerable to subsequent traumatic events. The discovery of lifelong GABA<sub>B</sub>-mediated intracortical inhibition alterations of the M1 gave rise to extensive ramifications into future clinical studies of sports concussions on associated functional losses and potential treatment alternatives.

One important discovery derived from alterations in intracortical inhibitory mechanisms in concussed athletes was inspired by pharmacological evidence suggesting that GABA<sub>B</sub> receptors play an important role in LTP<sup>60</sup> and learning.<sup>58</sup> Briefly, it is well known that synaptic plasticity occurring through LTP is an essential part of motor learning. In animal preparations, motor learning strengthens M1 synaptic efficacy and prevents subsequent LTP<sup>59</sup> from occurring without affecting synaptic modification range, which, according to the Bienenstock-Cooper-Munro theory of bidirectional synaptic plasticity,<sup>2</sup> is indicative of LTP involvement in motor skill learning. Recent advances in TMS research have enabled the noninvasive induction of LTP in human M1 to study its role in skill acquisition. In PAS paradigms, stimulation of the median nerve is repetitively paired with TMS of its homotopic representation in the M1, resulting in increased motor cortex excitability. Paired associative stimulation induction of LTP-like effects are reversible, topographically specific, and dependent on N-methyl-D-aspartate receptors.<sup>62,63</sup> Similar to what has been shown in animals, a period of motor learning prevents LTP-like plasticity from occurring after PAS, suggesting that motor learning occurs more easily in a system in which LTP induction is easily achieved. Knowing that the GABA<sub>B</sub> receptor agonist baclofen abolishes PAS-induced LTP-like plasticity, some have proposed that increases in GABA<sub>B</sub> receptor activity prevent LTP-dependent motor learning. Therefore, it stands to reason that excessive GABA<sub>B</sub> neurotransmission in the motor cortex of concussed athletes may reduce the probability of LTP induction, resulting in impaired motor learning. Recent findings consistent with this notion indicate that repeated concussions induced persistent elevations in GABA<sub>B</sub>-mediated intracortical inhibition in the M1, which was associated with suppressed PAS-induced LTP-like synaptic plasticity. This synaptic plasticity suppression was related to reduced motor learning relative to normal LTP-like synaptic plasticity in nonconcussed players. These findings reveal GABA neurotransmission alterations after repeated concussions and suggest that impaired learning in athletes with multiple concussions could at least partly be related to compromised GABA-dependent LTP synaptic plasticity.

**Motor Function in Late-Adulthood Athletes With Multiple Concussions**

Consistent with previous findings in young athletes who have suffered multiple concussions, TMS assessment of motor cortex excitability showed that the CSP was significantly prolonged in former athletes who had sustained their last sports concussion more than 3 decades prior to testing. Along with their prolonged CSP relative to controls, former athletes with a history of sports concussion exhibited significant slowness of movement, or bradykinesia, on an RAM task. Further correlational analyses established a strong relationship between the duration of the CSP and the movement velocity on the RAM task, such that former concussed athletes with more prolonged CSPs tended to be slower at executing alternating pronation-supination cycles. This finding suggests that the altered neurophysiological mechanisms that lengthen the CSP in concussed athletes could also be implicated in the biological bases of the slowness of movement seen
Enduring alterations in sports concussions

in former athletes with concussion when performing an RAM task.20

This significant slowness in executing RAM task cycles was found in former athletes with concussion who did not otherwise report experiencing motor difficulties in their daily activities. This finding is consistent with a recent study showing motor slowness on an RAM task in patients with very-early-stage Parkinson disease who had yet to experience the degenerative disease’s more debilitating symptoms.29 It remains to be verified in further longitudinal studies whether former concussed athletes who experienced early signs of movement slowness at the time of testing will proceed to incapacitating motor symptoms. Interestingly, motor execution speed on an identical RAM task was normal in young, active university football players with multiple concussions.30 The concept of cognitive reserve discussed earlier to explain the reappearance of neuropsychological test deficits in former athletes with concussions relative to nonconcussed counterparts could apply just as well to bradykinesia symptoms surfacing 3 decades postconcussion. In fact, the resilience of the young concussed brain in adapting to abnormal intracortical inhibition of the motor cortex appears to be sufficient to maintain normal motor execution speed, whereas the nature of an aging brain that had sustained prior concussions could no longer enable optimal performance levels.

Besides the depleted neural reserve, collected data cannot exclude the possibility that former concussed athletes were experiencing early symptoms of a neurodegenerative disease affecting motor system functions. This finding is in keeping with the catastrophic consequences of repetitive subconcussive and concussive head blows to athletes were experiencing early symptoms of a neurodegenerative condition.57 Among the cardinal motor impairments of chronic TBI, ataxia and motor execution slowness (bradykinesia) have both been related to significantly prolonged CSP duration in other neurodegenerative conditions. In fact, previous studies conducted in patients with cerebellar ataxia58 and Huntington disease40 have also revealed excessive CSP alongside bradykinesia/ataxia symptoms. Investigating the effects of chronic TBI relative to those of fewer remote concussions on the GABA receptor-mediated intracortical inhibition anomalies and associated motor symptoms could also help to establish the clinical significance of lifelong CSP anomalies after sports concussions.

Conclusions

In summary, the research to date has consistently shown enduring, cumulative cognitive and motor system function alterations after sports concussions that are present in both young concussed athletes and older former athletes who had previously sustained concussions. Strong converging evidence outlined in this review indicated that the damaging effects of concussion cannot and should not be overlooked any longer. Further, research efforts should be increased to deepen our understanding of the long-term pathophysiology of sports concussions and how it might be influenced by genetic and environmental factors. Additionally, investigating how more proactive injury management might influence the long-term and/or cumulative effects of the injury must be explored. Furthermore, we also highlighted the usefulness of more refined brain investigation tools to delineate the underlying causes and associated action mechanisms preventing the brain from fully recovering from the seemingly benign adverse effects of sports concussions, even if the persistent changes are subclinical. Indeed, the use of refined brain investigation tools is especially called for if we are to move from current management based mostly on expert opinion to an evidence-based medical approach. We also focused on the importance of implementing longitudinal follow-up studies of former concussed athletes, whose cognitive and motor system functional alterations may evolve into more severe, perhaps debilitating difficulties impairing daily function.

Disclosure

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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References

9. Cavanaugh JT, Guskiewicz KM, Stergiou N: A nonlinear dy-

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52. Omalu BI, DeKosky ST, Minster RL, Kamboh MI, Hamil-
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