Introduction

Strokes and movement disorders are quite common diseases and come to the attention of all physicians, but most often neurologists, emergency physicians, and primary care providers. Stroke-related (vascular) movement disorders, however, are relatively uncommon and represent a recognized complication of either ischemic or hemorrhagic strokes. Although the prevalence of post-stroke movement disorders is unclear because of the lack of prospective studies, data from retrospective studies have estimated the overall prevalence to range from 1% to 4% of all strokes with both sexes affected equally.1–3

The spectrum of post-stroke movement disorders is broad and includes both hypo- and hyperkinetic syndromes. The latter often present as variable combinations of hyperkinetic movements (dyskinesias) and can be broadly classified into three main phenotypes: choreiform dyskinesias (ballism, chorea, and athetosis), dystonia, and non-choreo-dystonic dyskinesias (eg, tremor, asterixis, and myoclonus) (Figure 1). The prevalence of vascular dyskinesias remains uncertain. In the Lausanne Stroke Registry from Switzerland, 29 (1%) of 2,500 patients admitted to the registry over 14 years developed hyperkinetic movement disorders with an estimated incidence of 0.08% per year. However, the large loss to follow-up has likely underestimated the prevalence, particularly for delayed movement disorders.1 In another stroke registry from Ecuador, 56 (3.7%) of 1,500 patients admitted to the registry over 9 years developed post-stroke movement disorders within the first year after stroke: 50 patients (3.3%) had hyperkinetic movements and 6 patients (0.4%) parkinsonism.2 Although data from these two studies suggest that vascular hyperkinetic movement disorders are likely to regress spontaneously, it remains unknown whether that decreases their...
prevalence over time as the incidence of delayed vascular
dyskinesias is largely unidentified.

This review summarizes the current knowledge about
the complex pathophysiology, overlapping phenomenology,
and treatment options for common post-stroke hyperkinetic
movement disorders. We searched for related published
literature via PubMed through June 2016 by using the
search terms “stroke”, “vascular”, “dyskinesia”, “movement
disorders”, “basal ganglia circuit”, “cerebellar circuit”, and
“motor control”.

Terminology
Stroke is broadly defined as an episode of neurological
dysfunction caused either by a focal infarction of the central
nervous system (CNS) or by a nontraumatic intracerebral
or subarachnoid hemorrhage.4 Although acute focal neuro-
logical deficits (eg, hemiparesis) are the clinical hallmark
of stroke presentation, nonfocal deficits (eg, impaired
consciousness) or positive neurological phenomena
(eg, hemidyskinesia) can occasionally be the initial mani-
festation of stroke resulting in diagnostic uncertainty and
delayed treatment.4-6

Movement disorders are a group of basal ganglia and/or
cerebellar circuit disorders characterized by impaired ability
to control or coordinate movements. They are usually clas-
sified first based on an accurate description of the abnormal
movement (phenomenology), and then subdivided according
to the underlying cause (etioloogy). The “phenomenological
classification” divides movement disorders into the following
types: 1) Hypokinetic (parkinsonian) disorders, dominated by
poverty (hypokinesia) or slowness (bradykinesia) of move-
ment, 2) Hyperkinetic (dyskinetic) disorders, characterized
by excessive, abnormal involuntary movements, and 3) Other
movement disorder syndromes, cannot easily be grouped
under the previous two categories, such as ataxia and akath-
isia. Furthermore, hyperkinesias (dyskinesias) are classified
as “focal” if only one body region is involved, “segmental”
if ≥2 adjacent body regions are affected, “multifocal” if ≥2
noncontiguous body regions are affected, “hemibody” if the
ipsilateral arm and leg are involved, and “generalized” if the
trunk and ≥2 other body regions are involved. The “etio-
logical classification” of movement disorders, on the other
hand, subdivides them into the following types: 1) Primary
(genetic or idiopathic) disorders, without an identifiable
secondary cause; 2) Secondary (symptomatic) disorders,
due to a known acquired etiology such as vascular, toxic, or
metabolic abnormalities; and 3) Psychogenic (functional)
disorders, commonly due to conversion, somatic symptom,
or factitious disorders.7

Although “dyskinesia” is a broad term that can indicate
any hyperkinesia, it is often employed in clinical practice
to indicate mixed, unusual, or complex hyperkinetic move-
ments, especially when these movements are challenging to
describe; nevertheless, a more specific phenomenological
categorization should be used whenever possible. For
example, variable mixtures of ballism, chorea, and athetosis
are often seen simultaneously in the same patient and are
commonly referred to as “choreiform dyskinesias”. Not
infrequently, dystonia coexists with choreiform movements
and the term “choreo-dystonic dyskinesia” is often used in
this clinical setting. Interestingly, the choreiform dyskine-
sias are considered variants within the same phenotypic
spectrum, with ballism and athetosis representing its fastest
and slowest motor phenotypes, respectively (Table 1).7-11
Finally, the term “dyskinesia” is also used more specifically
to indicate certain movement disorders such as levodopa-
induced dyskinesia (due to chronic exposure to levodopa in
Parkinson’s disease patients) and tardive dyskinesia (due to
prolonged exposure to dopamine-blocking agents).12,13

Pathophysiology
Motor control is a complex process that is governed by
a sophisticated motor circuitry involving both pyramidal
(cortical) and extrapyramidal (basal ganglionic and cerebellar)
circuits. Motor commands are generated in the motor cortex,
but basal ganglia and cerebellum closely refine these signals by
acting as feedback loops to allow for smooth, accurate, coor-dinated
movements. While the cortico-basal loop (basal ganglia
circuitry) provides a tonic inhibitory output to the thalamus and
thus the motor cortex, the output of the cortico-cerebellar loop
(cerebellar circuitry) is tonically excitatory. Glutamate is the
Table 1 Major phenomenological types (phenotypes) of hyperkinetic movement disorders

<table>
<thead>
<tr>
<th>Dyskinesia</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Ballism (Ballismus)</td>
<td>Violent, irregular, large-amplitude, involuntary movements that mainly involve the proximal extremities. It can be viewed as a fast form of chorea.</td>
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<tr>
<td>Chorea</td>
<td>Rapid, irregular, involuntary movements that typically involve both proximal and distal muscles.</td>
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<tr>
<td>Athetosis</td>
<td>Slow, irregular, writhing, involuntary movements that mostly involve the distal extremities (hands or feet). It can be viewed as a slow form of chorea.</td>
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<tr>
<td>Dystonia</td>
<td>Sustained or intermittent involuntary muscle contractions causing abnormal postures and/or repetitive movements of the affected body part.</td>
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<tr>
<td>Tremor</td>
<td>Rhythmic, involuntary oscillations of a body part. Tremors are broadly classified based on phenomenology into rest and action (postural and/or kinetic) tremors. Kinetic tremors are then subdivided into simple (present throughout the movement), intention (increasing throughout the movement), or terminal (occurring at the end of the movement).</td>
</tr>
<tr>
<td>Myoclonus and asterixis</td>
<td>Sudden, brief, jerky, arrhythmic, involuntary movements caused by muscle contractions (myoclonus = positive myoclonus) or loss of muscle tone (asterixis = negative myoclonus).</td>
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</table>

Major excitatory neurotransmitter and gamma-aminobutyric acid (GABA) is the major inhibitory neurotransmitter in this motor network (Figure 2). Despite the significant overlap in their functions, the basal ganglia are particularly important for gross motor skills, whereas the cerebellum is crucial for fine or complex motor skills (eg, speech and writing) in which accurate performance is necessary.

The basal ganglia are a large collection of subcortical nuclei, which play a central role in modulating motor cortical activity through the selection and execution of appropriate motor programs. Almost all inputs to the basal ganglia circuitry arrive via the striatum (caudate and putamen). The motor signals are then processed through the pallidum (globus pallidus interna “GPi” and externa “GPe”), motor thalamus (ventral anterior “VA” and ventral lateral “VL” nuclei), and then back to the motor cortex. The normal function of this motor loop is influenced by two additional nuclei: substantia nigra and subthalamic nucleus, which regulate the activity of the striatum and pallidum, respectively. While glutamate and GABA are the major excitatory and inhibitory neurotransmitters, respectively, dopamine and acetylcholine act as neuromodulators in this circuit that regulate striatal activity. Dopamine is the transmitter of the nigrostriatal pathway, whereas acetylcholine is a major transmitter of striatal interneurons (Figure 3).

The striatum is the largest component of the basal ganglia motor circuit, and the vast majority of its neurons are GABAergic medium-sized spiny neurons whose axons comprise the striatal output pathways. These neurons are divided into two subsets of approximately equal numbers and provide two projection systems that have opposite effects on movement, the direct and indirect pathways. The direct pathway (GO pathway) originates from dopamine D1 receptor-expressing neurons and projects directly to the GPi. It is an excitatory pathway because its net effect is.

![Figure 2](https://www.dovepress.com/)

Figure 2 A simplified diagram of the motor control circuitry. The motor cortex projects to the cerebellum and basal ganglia and receives feedback signals from them. Although cerebellar output is tonically excitatory and basal ganglia output is tonically inhibitory, the balance between these two systems is of pivotal importance for motor control and coordination. GLU and GABA are the major excitatory and inhibitory neurotransmitters in this network, respectively.

Abbreviations: GLU, glutamate; GABA, gamma-aminobutyric acid.

![Figure 3](https://www.dovepress.com/)

Figure 3 A simplified diagram of the basal ganglia motor circuitry, including its direct and indirect pathways. DA plays a central role in this circuit and acts as a neuromodulator that regulates the striatal function.

Abbreviations: DA, dopamine; GLU, glutamate; GABA, gamma-aminobutyric acid; GPe, external globus pallidus; GPi, internal globus pallidus.
to disinhibit the thalamus and thus the motor cortex. The indirect pathway (NO-GO pathway) originates from D2 receptor-expressing neurons and projects to the GPe and subthalamic nucleus before terminating in the GPi. It is an inhibitory pathway because its net effect is to inhibit the thalamus and thus the motor cortex. Dopamine has differential effects on these two pathways: it activates D1 receptors and facilitates the direct pathway, but it inhibits D2 receptors and suppresses the indirect pathway. Therefore, the net effect of dopamine is to facilitate voluntary movements by promoting transient interruptions of the tonic inhibitory output of the basal ganglia via the simultaneous activation and suppression of the direct and indirect pathways, respectively.16–20

The cerebellum consists of a cortex, white matter, and deep cerebellar nuclei. It acts as a sensorimotor information processor by receiving information from all parts of the nervous system and comparing the motor commands of the cortex with the proprioceptive information coming from joints and muscles. This enables the cerebellum to detect errors in muscle contractions during active movements and thus to contribute to motor accuracy and coordination.15

Two cerebellar feedback loops are critical for this task, the dentato-rubro-olivary loop and the cortico-cerebellar loop. The dentato-rubro-olivary circuit (Guillain-Mollaret triangle) connects the dentate nucleus in the cerebellum with the contralateral red nucleus and inferior olivary nucleus in the brainstem via the superior cerebellar peduncle, the central tegmental tract, and the inferior cerebellar peduncle, respectively.21 This subcortical circuit is itself part of the larger cerebellar motor network, the cortico-cerebellar circuitry, in which the cerebral and cerebellar cortices are connected together indirectly. The motor cortex projects to the cerebellar cortex (via pontine nuclei) which, in turn, projects primarily to the VL nuclei of thalamus and then back to the cerebral cortex (Figure 4).14,15,23 As the inferior olivary nucleus receives collateral inputs from all afferent pathways projecting to the cerebellar cortex via mossy fibers, it compares intended with executed movements and conveys error signals to the cerebellar cortex via climbing fibers.14,15,24

The pathogenesis of stroke-related dyskinesias is still incompletely understood but suggested mechanisms include post-synaptic denervation hypersensitivity, trans-synaptic neuronal degeneration, as well as aberrant axonal and dendritic plasticity (remodeling) after the cerebrovascular injury.3 Post-stroke dyskinesias are relatively rare even with marked lesions, but they can arise after any stroke subtype at any level within the motor circuitry and after any interval period. Furthermore, no specific anatomical locations in the motor circuitry are reliably predictive of a particular dyskinesia, whereas the same dyskinesia can be caused by lesions in different locations in this circuitry. Due to this overlap, stroke-related movement disorders cannot be predicted from the location, size, or number of vascular insults.1,2,19 Interestingly, though the pathogenesis of vascular and primary movement disorders is different, they share similar underlying pathophysiology.

If symptomatic, vascular lesions involving the basal ganglia circuitry usually present with contralateral abnormal involuntary movements, but there have been rare case reports of ipsilateral dyskinesias.25 The most commonly involved areas in this regard are the striatum followed by the thalamus, but other reported locations include the fronto-parietal cortex, caudate, subthalamic nucleus, corona radiata, internal capsule, and pons.1,3,19 These vascular dyskinesias are believed to arise from underactivity of the indirect pathway (and/or overactivity of the direct pathway) leading to a decreased pallidal inhibitory output to the thalamus. The resultant thalamic disinhibition releases the motor cortex and allows movements that are normally suppressed.2,19,26,27

Vascular parkinsonism, on the other hand, is usually caused by diffuse or multiple insults in the basal ganglia loops and presents with non-tremulous, lower-body parkinsonism.
(ie, predominantly affecting the legs). It is believed to arise from overactivity of the indirect pathway (and/or underactivity of the direct pathway) leading to an increased pallidal inhibitory output to the thalamus. The resultant thalamic inhibition suppresses the motor cortex and leads to hypokinetic movements.\textsuperscript{3,23,29}

The cortico-cerebellar and dentato-rubro-olivary circuits are critically involved in tremor genesis.\textsuperscript{30} Insults to the various structures of this network can give rise to cerebellar tremor, palatal tremor (previously known as palatal myoclonus), or Holmes tremor (also known as thalamic, rubral, midbrain, or cerebral outflow tremor). The latter occurs when the lesion involves both the nigrostriatal and cerebellar pathways resulting in resting and action tremor, respectively.\textsuperscript{31–33} Dystonic and parkinsonian tremors are special forms of tremor that occur in association with vascular dyskinesia and parkinsonism, respectively.

Differential Diagnosis

Acute focal, segmental, or hemibody dyskinesia should always raise a suspicion of structural lesions involving the motor circuitry.\textsuperscript{11,33} Although stroke is the most common cause after the age of 50 years, other structural pathologies have also been reported, including tumors, arteriovenous malformation, cerebral abscess, encephalitis, and multiple sclerosis.\textsuperscript{11,33–36} Non-structural causes, especially hyperglycemia and nonketotic hyperglycemia, can occasionally present with dyskinesia.\textsuperscript{37–41} Interestingly, hemichorea-hemiballism is the most commonly reported dyskinesia arising secondary to either structural (eg, vascular) or non-structural (eg, dysglycemic) etiologies.\textsuperscript{34,36,38,41}

Dyskinesias, but isolated dyskinesia unaccompanied by other abnormal movements is rare. Action tremor is by far the most common form, but rest tremor can occasionally be seen (eg, in vascular parkinsonism). Cerebellar tremor, the prototype vascular tremor, is primarily a slow intention tremor (frequency < 5 Hz) with frequent postural component. Holmes tremor is usually a low-frequency tremor (< 5 Hz) with rest, postural, and intention components in the affected upper extremity. Palatal tremor is a slow tremor of the soft palate (< 5 Hz) in which patients may complain of a disturbing clicking sound generated by palatal muscle contractions causing opening and closing of the eustachian tube. Dystonic tremor is usually a focal postural and/or kinetic tremor that occurs in association with dystonia, which might be subtle or overshadowed by the tremor. Despite being arrhythmic with variable frequencies, dystonic oscillations are traditionally referred to as “tremor.”\textsuperscript{50}

Post-stroke myoclonus and asterixis are usually focal or segmental with corresponding lesions reported in numerous, mostly contralateral, brain regions. Myoclonus is not an uncommon component of mixed vascular movement disorders, but isolated myoclonus is rare. Dystonic myoclonus
(myoclonic dystonia) has been described in patients with vascular dystonia secondary to thalamic infarcts.2,3,19

Prognosis

The natural history of post-stroke dyskinesias is variable. Their onset can be either “early/acute”, occurring shortly after stroke, or “delayed/chronic”, emerging months to years later. On the other hand, their course can be transient, recurrent, persistent, or progressive.1,43,44 Although the latency interval seems to depend partially on the dyskinesia type, it still varies widely within each vascular hyperkinetic movement disorder.2,3 While dystonia is frequently delayed in onset, hemichorea-hemiballism usually occurs shortly after stroke and occasionally represents the initial manifestation of acute cerebral ischemia or intracerebral hemorrhage.2,3,33,45,51 Post-stroke dyskinesias are usually self-limited and resolve within 6 to 12 months of onset, but the overall long-term prognosis of the affected patients is similar to that of other stroke patients.1,2,46,52

Management

Although post-stroke dyskinesias tend to resolve spontaneously, a short-term treatment might sometimes be required for symptom control.1,2 As in all cases of secondary movement disorders, treatment of the underlying etiology is of paramount importance. Control of vascular risk factors is crucial in reducing the incidence of vascular dyskinesias. The discussion of stroke prevention and management, however, is beyond the scope of this article. Symptomatic pharmacotherapy might be necessary for severe dyskinesias, but periodic trials of therapeutic withdrawal (ie, for patients with controlled symptoms) are required due to the high likelihood of spontaneous regression. Medications should be started at low doses and gradually titrated up until an effective and tolerable dosage is reached. Although there are no established treatment guidelines, most of the treatment options are similar to those for primary movement disorders based on similar underlying pathophysiology.

The symptomatic pharmacotherapy for post-stroke choreiform dyskinesias consists mainly of anti-dopaminergic therapy with typical or atypical antipsychotics (neuroleptics). Blockade of striatal dopamine D2 receptors is believed to be responsible for their anti-dyskinetic activity, as D2 receptor antagonists disinhibit the indirect pathway and, therefore, suppress abnormal involuntary movements. Unfortunately, dopaminergic blockade carries the risk of acute dystonic reactions, tardive dyskinesia, and drug-induced parkinsonism. Atypical antipsychotics (eg, risperidone), however, are less likely to cause these side effects compared to typical antipsychotics (eg, haloperidol) and, therefore, are generally preferred for this use.53,54 Tetrabenazine, a presynaptic dopamine depletor with weak postsynaptic D2 receptor blocking activity, is a reasonable alternative for patients who are intolerant or unresponsive to dopamine receptor antagonists.55 Tetrabenazine acts primarily as a reversible inhibitor of the vesicular monoamine transporter 2 (VMAT2) in the presynaptic nerve terminals, thus exposing dopamine and other monoamines to monoamine oxidase and leading to their depletion.56 Unlike dopamine receptor blockers, tetrabenazine rarely causes tardive dyskinesia because of its dominant presynaptic anti-dopaminergic properties.3,37 Non-dopaminergic drugs have been tried in the management of vascular choreiform dyskinesias with varying success. Case reports and small case series have suggested a potential beneficial effect of antiepileptic drugs including levetiracetam, topiramate, gabapentin, clonazepam, and valproate.58-62

Chemodenervation with botulinum toxin (BTX) injections is the cornerstone of the symptomatic treatment of focal or segmental post-stroke dystonia.1 BTX is a neurotoxic protein with several serotypes that cleaves the synaptic proteins (SNARE) in the presynaptic nerve terminals, thereby blocking the release of acetylcholine at the neuromuscular junction. SNARE proteins are required for the fusion of presynaptic storage vesicles containing acetylcholine with the presynaptic membrane.53 BTX injections are given intramuscularly, often under electromyography (EMG) guidance, and need to be repeated every 3 to 6 months.64 Vascular dystonia usually has a poor response to oral pharmacotherapy, including dopamine blocking and depleting agents, anticholinergic drugs, baclofen, and benzodiazepines. Oral medications, however, are widely used in generalized dystonia, dystonia mixed with other movement disorders, or as an adjuvant therapy in focal or segmental dystonia when there is an unsatisfactory response to BTX.64-67

Post-stroke tremor is particularly refractory to pharmacotherapy. Trials of medications with GABA-agonistic activity (eg, clonazepam, valproate, topiramate, or primidone), alone or in combination, may be effective in individual cases. Because of the dopaminergic (nigrostriatal) system involvement in Holmes tremor, treatment with levodopa or dopamine agonists seems to be useful.31 Propranolol, one of the first-line treatments in essential tremor, is usually of limited benefit in vascular tremor. Dystonic and parkinsonian tremors are treated as vascular dystonia and parkinsonism, respectively.3,19

Post-stroke myoclonus and asterixis usually improve spontaneously and do not require pharmacotherapy. When interfering with the patient’s functional abilities, like eating or writing, myoclonus is most frequently treated with
GABAergic medications (eg, clonazepam and valproate), but levetiracetam or piracetam can be very useful. Monotherapy should be attempted first, although eventually several drug combinations might be required. Post-stroke dyskinesias is treated as vascular dystonia.

Finally, stereotactic functional neurosurgery, whether ablative or deep brain stimulation (DBS), should be considered for patients with severe and persistent dyskinesias (arbitrarily defined as duration longer than 1 year). While lesioning procedures (eg, pallidotomy or thalamotomy) are carried out using radiofrequency ablation or gamma knife radiosurgery, DBS (eg, pallidal or thalamic DBS) uses high-frequency electrical stimulation of the targeted nuclei. Although the therapeutic mechanisms of action are not completely understood, ablative surgeries are believed to destroy abnormally hyperactive circuits in deep brain nuclei, while DBS suppresses abnormally excessive activity in the motor circuitry without significant destruction of the brain tissue. Neurostimulation, therefore, has largely replaced neuroablation to treat refractory dyskinesias because it is less invasive, adjustable for maximal symptomatic benefit, and reversible in case of adverse effects. Stereotactic functional neurosurgery depends on the accurate identification of anatomically and functionally distinct deep brain structures to maximize therapeutic benefits and to minimize adverse neurological events of any surgical intervention. The contralateral motor thalamus or internal pallidum are the usual neurosurgical targets, but it is yet to be determined which target is more effective for specific dyskinesias. The ventrolateral “VL” nuclei, including ventralis intermedia and ventralis oralis posterior, are the most common nuclei targeted within the motor thalamus. Potential candidates with vascular dystonia should undergo surgery prior to the development of contractures and fixed deformities that may limit functional improvement as the dystonia improves.

Conclusion
Stroke is the leading cause of focal or segmental limb dyskinesias as well as hemidyskinesia in elderly patients, but other structural and non-structural brain abnormalities should also be considered in the differential diagnosis. Post-stroke dyskinesias, which can arise from either ischemic or hemorrhagic cerebrovascular insults, are often mixed and variable with several components of hyperkinetic movements. A comprehensive clinical and neuroimaging assessment is essential to establish the correct diagnosis. Fortunately, post-stroke dyskinesias tend to resolve spontaneously within 6 to 12 months. A short-term symptomatic pharmacotherapy may be required in some patients, whereas surgical treatment is reserved for persistent, disabling, and medically intractable cases.

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