Pharmacological Aspects of Neurorehabilitation

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ABSTRACT
Physicians in neurorehabilitation often deal with pharmacological problems, marshalling antihypertensive, anticonvulsive and antiocoagulation treatments. In addition, there is growing interest in positive or negative effects of medication on brain recovery. Of great importance is the concept of so-called “detrimental drugs” known to negatively influence processes of brain reorganization and recovery. To this group belong anti-convulsive agents such as phenytoin and barbiturates as well as benzodiazepines, butyrophenones and the antihypertensives clonidine and prazosine. Whenever possible these drugs should be avoided in the course of brain recovery after a cerebral lesion.

For only two substances (the SSRI fluoxetine and cerebrolysin, a mixture of pleotropic neuropeptides and amino acids) large randomized controlled trials showed a positive influence on facilitating motor recovery after the stroke. Both substances probably work through pleotropic multiple molecular mechanisms and not as a one-to-one agonist on the receptor. In general the use of antidepressive agents especially SSRI after the stroke can also be recommended for non-depressed stroke patients.

Also dopaminergic drugs have been shown in smaller studies to positively influence functional recovery. Considering their low side-effect profile, the tentative use of 100 mg of L-Dopa per day in the subacute phase of the stroke can be recommended. In MS patients the use of antidepressive agents is also recommend to improve life quality.

In patients with diminished states of consciousness amantadine is the only substance which a randomized controlled study proved to have at least some transient effect. The use of amantadine can be recommended for the improvement of the level of consciousness in these patients.

Introduction
Physicians engaged in neurological rehabilitation constantly have to deal with aspects of primary pharmacological treatment of patients, including control of high blood pressure, anticonvulsive therapies and suitable antiocoagulation treatment to reduce risk factors and secondary problems. Furthermore, neurological rehabilitation must also take into account pharmacological issues relating to restoration of brain function. This concerns the avoidance of pharmaceuticals that may interfere with brain recovery as well as the use of drugs that may have a positive affect on brain function. This overview aims to provide a critical summary of the options available to the clinician in the pharmacological treatment of patients after acute neurological events as part of the process of the rehabilitation of brain organization and restoration of brain function, as well as discuss the avoidance of potentially negative effects of pharmacological interventions.

General pharmacology of restoration of brain function
There are good animal-experimental data for numerous commonly-used drugs regarding their influence of spontaneous brain recovery after acute neuronal damage.

These data were developed based on experimental trauma or ischemia models. The models [1–5] convincingly demonstrated that as a rule amphetamines significantly improved motor and perceptual functions [6–8]. Amphetamines demonstrate an inverse U-shaped relationship with an optimal effect at medium doses, and their use depends on parallel, task-related behavior. As a rule they are blocked by haloperidol [9–11].

The likely action mechanism of action of amphetamines is the central release of norepinephrine [12, 13].
antagonize the effect of amphetamines [9, 10, 16, 17]. In contrast, intrusion of spontaneous brain recovery in all experiments and can brain function restoration [14–15].

Norepinephrine + +
Amphetamine + +
Clonidine − −
Prazosin − ?
Haloperidol − −
Propanolol 0 −
GABA − −
Diazepam − −
Muscimol − −
Phenytoin − ?
ACH + +
Scopolamine − −

*: Promoted effect 0: No effect −: Reduced effect

Other sympathomimetic substances such as methylphenidate and alpha-2 antagonists including yohimbine and idazoxane also increase norepinephrine release and lead to an improvement in brain function recovery. Conversely, alpha-2 agonists such as clonidine or alpha-1 antagonists such as prazosin may interfere with brain function restoration [14–15].

Haloperidol and other butyrophenones show a considerable disruption of spontaneous brain recovery in all experiments and can antagonize the effect of amphetamines [9, 10, 16, 17]. In contrast, classical tricyclic antidepressants such as clomipramine or imipramine show only neutral or slightly negative influences on brain recovery.

In animal experiments, serotonin reuptake inhibitors (SSRI) such as trazodone or fluoxetine have neutral or slightly positive effects on brain function recovery [18]. Significantly negative effects can be described for most GABAergic substances, especially for benzodiazepines [19–21].

The same applies to many classical anticonvulsant substances such as phenytoin and barbiturates [22, 23]. It has long been known that they can adversely affect spontaneous brain recovery, probably via GABAergic mechanisms [24, 25]. After experimental infarctions, anticholinergics have also been shown to disrupt motor recovery. On the other hand, carbamazepine does not appear to influence recovery mechanisms.

It is not entirely clear which molecular mechanisms in the indicated substances are responsible for promoting or inhibiting restoration of brain function. It is highly likely that influences on long-term potential (LTP) play an essential role in this regard. As shown in Table 1, there is a clear parallel between the effects of some substances influencing the brain recovery function and their effect on LTP as demonstrated in animal experiments [26].

### The “detrimental drug” concept

Many of the above-mentioned pharmacologically active brain function-enhancing drugs are still uncritically used in everyday clinical situations in patients after suffering severe cerebral lesions.

Larry Goldstein was the first to use introduce the concept of so-called detrimental drugs [27].

In catamnestic studies he showed that after brain injury, patients receiving one of the substances listed in Table 1 showed significantly worse functional end results after months or years of treatment compared to patients who were not treated with any substances which might potentially damage brain organization. It cannot be ruled out with certainty that these effects are based on selection bias, since it is possible that the patients treated with detrimental drugs had more severe brain damage.

The data on the use of detrimental drugs is, of course, inconsistent, and recommendations are based essentially on animal-experimental data. Randomized controlled studies are lacking completely and are probably not ethically justified as well.

Nevertheless, it is advisable to dispense with these substance classes in clinical practice in the acute and post-acute phase after stroke, cerebral trauma and other brain injury. Today it is highly likely that ruling out detrimental drugs is clinically more important than administration of substances promoting brain reorganization.

### Substances improving restoration of brain function

#### Amphetamines

Various substances influencing monoaminergic transmission have been used experimentally to improve brain function. After the early animal experiments of Feeney and colleagues [9], there was great interest in introducing amphetamines into clinical practice in order to positively influence brain function recovery procedures. This has been supported by several experimental studies in humans showing that amphetamines increase the capacity of motor learning [28] and, for example, sensitive two-point discrimination in normal subjects [29].

In 1988, Chrisostomo et al. [30] performed an initial study of only 8 patients which showed some positive effects of amphetamine on patients in the subacute phase post-stroke.

Larger studies have shown varying and unconvincing results [31–42]. Therefore, many open questions remain regarding the use of amphetamines, e.g. dosage at which the amphetamine com-

**Table 1 Parallel effects of individual substances on function recovery or long-term potential (LTP).**

<table>
<thead>
<tr>
<th>Substance</th>
<th>Effect on function recovery</th>
<th>Effect on LTP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Norepinephrine</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Amphetamine</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Clonidine</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Prazosin</td>
<td>− ?</td>
<td></td>
</tr>
<tr>
<td>Haloperidol</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Propanolol</td>
<td>0</td>
<td>−</td>
</tr>
<tr>
<td>GABA</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Diazepam</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Muscimol</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>− ?</td>
<td></td>
</tr>
<tr>
<td>ACH</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Scopolamine</td>
<td>−</td>
<td>−</td>
</tr>
</tbody>
</table>

*: Promoted effect 0: No effect −: Reduced effect

**Table 2 List of so-called detrimental drugs (substances inhibiting restoration of brain function).**

<table>
<thead>
<tr>
<th>Detrimental drugs</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haloperidol</td>
<td>D2 antagonist</td>
</tr>
<tr>
<td>Prazosin</td>
<td>Alpha-1 antagonist</td>
</tr>
<tr>
<td>Clonidine</td>
<td>Alpha-2 agonist</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>GABA mechanism</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>GABA agonist</td>
</tr>
<tr>
<td>Phenobarbital</td>
<td>GABA mechanism</td>
</tr>
</tbody>
</table>
ponents should be used, at which time and for what duration in the acute or subacute or chronic phase after stroke. Therefore, the question of the usefulness of amphetamines cannot be conclusively clarified, and there is no recommendation regarding amphetamines during the subacute or chronic phase after stroke.

At the moment it can be said that although the use of amphetamines has some potential to improve brain function recovery after stroke, clinical use cannot be recommended with respect to motor or cognitive functions.

**Dopaminergics**

Levodopa is a pharmacologically “dirty” substance because it affects not only dopaminergic but also partially influences noradrenergic neurotransmission. A first randomized controlled study by Scheidtmann et al. [43] compared 22 stroke patients treated with levodopa with 25 receiving a placebo; the results was that using levodopa resulted in an increase in motor function (dose: 100 mg L-Dopa for 6 weeks after onset).

It is unclear to what extent the efficacy of dopaminergics depends on the duration of the dose after an acute lesion. Perhaps longer-term administration of a few weeks after an acute event is more promising than a too short administration of the drugs. Of course, a final clarification of this issue is only possible through systematic prospective randomized trials.

This finding of dopaminergics could not be confirmed in subacute to chronic stroke patients who received L-Dopa for 9 weeks [44]. It was demonstrated that L-Dopa improved the learning of word lists by acute stroke patients compared to those receiving a placebo [45]. Likewise, Senlow et al. [46] showed that patients receiving L-Dopa prior to the start of speech therapy showed improved speech recovery (word fluency and repetition of word lists). On the other hand, Leeman et al. [47], in a crossover trial of subacute administration of L-Dopa for 2 weeks indicated no positive effects on speech recovery.

Restemeyer et al. [48] observed no significant effect of a single dose of L-Dopa on motor recovery and motor excitability in chronic stroke patients. In a long-term administration of L-Dopa in chronic stroke patients, Lokk et al. [49] showed that four-week administration of L-Dopa with a single daily dose improved motor performance. Similar results as in the Scheidtmann et al. study were confirmed in a study by Masihuzzaman et al. [50]. In a case control design, stroke patients were either randomized to physiotherapy in combination with L-Dopa or physiotherapy alone with placebo for 8 weeks. The L-Dopa group showed greater improvement in the Rivermead Mobility Index.

Further studies [51–54] have investigated the outcome of L-Dopa in combination with physiotherapy in chronic stroke patients years after the stroke event. Various study designs have shown somewhat positive effects. The 2005 study by Floel et al. [53] demonstrated positive effects using transcranial magnetic stimulation on indicated potentials but no clinical effects. Restemeyer’s study [48] indicated no positive clinical effects while other studies with different designs reflected positive effects on motor learning and various motor scores [51, 52, 54]. It should be emphasized that all these studies of chronic patients were based on very small samples (only up to 18 patients).

There are currently no studies of dopaminergic facilitation using robot-assisted therapy or virtual reality. Recently Tran et al. [55] published a positive review of the utilization of dopaminergics.

In summary it can be stated that there is some evidence that positive results can be achieved by daily dosages of 100 mg of L-Dopa in both subacute and chronic stroke patients if administered over the course of at least one week. A final decision on effectiveness cannot be made at this time, however, and thus there is no clear recommendation to use L-Dopa. However, in view of the low potential for side effects, administering L-Dopa can be useful.

**Reboxetine**

Reboxetine is a selective norepinephrine reuptake inhibitor approved for treating depression. Based on its pharmacological profile it can be assumed that this substance has properties similar to amphetamines in the facilitation of noradrenergic neurotransmission. Considering that reboxetine has very few side effects compared to amphetamines (no addiction potential, little pulmonary risk), this substance can be regarded as useful for improving brain function recovery. Initial investigations showed positive effects on restoration of motor-related brain function [56, 57].

**Methylphenidate**

Methylphenidate is a commonly used psycho-stimulant in the treatment of attention deficit disorder (ADD)/hyperactivity problems. This is a piperidine component that increases the concentration of dopamine and norepinephrine by inhibiting multiple monomodal transporters.

Numerous studies have investigated a possible effect of methylphenidate on cognitive recovery after stroke and brain trauma [58–60]. Early administration of methylphenidate after severe cerebral trauma could shorten the length of stay in the intensive care unit [61].

Most of the studies with methylphenidate were limited to very small groups of patients, so that definite conclusions on its fundamental efficacy cannot currently be made.

**Amantadine**

Amantadine is a substance with various pharmacological characteristics, such as increasing the release of striatal dopamine, retarding dopamine reuptake while increasing the number of postsynaptic dopamine receptors. Furthermore, amantadine stimulates DOPA decarboxylase in the striatum while acting as a weak noncompetitive NMDA glutamate receptor antagonist similar to memantine, often used in the treatment of degenerative dementia [62].

Amantadine was first introduced as an antiviral substance, especially against influenza viruses [63]; it was subsequently discovered that amantadine also had a positive effect on parkinsonian symptoms [64].

Smaller experimental studies suggested a positive effect of amantadine on improving states of restricted consciousness in cerebral trauma [65–70].

In an open study, Kraus et al. [67] demonstrated a significant improvement of executive functions by administering amantadine
to SHT patients, combined with a distinct increase in left-hemispheric prefrontal glucose metabolism in PET studies.

Giacino et al. [71] in a methodically high-value placebo-controlled double-blind, multi-center investigation of 184 patients with various pronounced states of limited consciousness (non-responsive and minimally-responsive wakefulness state) demonstrated that in comparison to the placebo group, recovery of patients receiving amantadine during a 4-week interval was accelerated. After amantadine was discontinued, however, the recovery slowed down so that after a total of 6 weeks, both groups exhibited similar results. Likewise, irritability after SHT is positively influenced by amantadine [72]. It is also currently too early to derive definitive conclusions regarding this substance. The large study by Giacino [71] did show, however, that amantadine has a demonstrable therapeutic potential, particularly for patients in a state of limited consciousness.

Piracetam

Piracetam is a nootropic substance frequently used to treat dementia, and chemically is a derivative of γ-aminobutyric acid. With respect to neurorehabilitation, this substance is interesting beyond treating dementia, since several randomized controlled studies have demonstrated its effect in the treatment of patients with severe aphasia [73–75]. This has likewise been confirmed by a Cochrane review [76].

The exact mode of action is unclear, but its effect on AMPA and NMDA glutamate receptors is likely. It is not possible to derive reliable dosage data based on the literature, since dosages varied from 4.8 g/day [73, 77] to 12 g/day [78].

From the clinical point of view, it is certainly helpful to use the substance in high intravenous doses (12 g per day) in the acute/subacute phase in patients with severe to moderate aphasia.

Anti-depressants

Depression is a common secondary problem in patients after stroke and cranial trauma which can significantly affect their cognitive and motor abilities [79].

The frequency of depression after stroke has been indicated to be 60 % or more [80]. In the mid-1980s, Reding was the first [81] in the USA to show that treating depression in stroke patients not only improved their depression, but also had a positive effect on the rehabilitation process. Several placebo-controlled studies have shown that the administration of antidepressants such as trazodone resulted in significantly better function recovery measured by relevant scores and also resulted in an improvement of ADL activities compared to using a placebo.

Dam et al. [82] demonstrated that the use of serotonin reuptake inhibitors such as fluoxetine and maprotiline resulted in improved motor recovery in stroke patients. Early use of antidepressants particularly resulted in improvement of cognition [83]. In addition, early treatment with antidepressants after stroke (within the first 12 weeks) showed a significant increase in life expectancy [84]. This applies to depressive and non-depressive patients alike. The mechanisms contributing to this improvement in life expectancy resulting from early treatment with antidepressants are unclear.

Chollet and colleagues [85] then performed the best evidence-based study of motor recovery after acute stroke according to EBM criteria. Their FLAME trial treated a total of 118 patients suffering from severe motor deficits with fluoxetine (20 mg daily) or a placebo for 3 months, starting 5–10 days post-stroke. Twenty days after treatment, motor scores showed an improved recovery of function using fluoxetine as compared to placebo.

Multiple mechanisms contribute to the positive effect of antidepressant drugs on motor and cognitive recovery after stroke.

Improving their mood can result in patients being more willing and motivated to participate in rehabilitation therapies. In addition, there is probably a more complex influence of antidepressants on brain function recovery which goes beyond pure 1:1 transmitter-receptor interaction. It has been known for many years that antidepressants also develop their “classic” antidepressant effect only weeks after the start of the treatment. The cause is certainly a complex intervention of the antidepressants in the mechanisms of brain reorganization, and therefore not only in the stroke treatment itself, but also with respect to normal endogenous depression. Manifold molecular mechanisms play a role in this (see Table 3).

### Table 3 List of antidepressive effects on various monaminergic transmitter systems.

<table>
<thead>
<tr>
<th>Acute Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>NA reuptake inhibitor</td>
</tr>
<tr>
<td>5-HT reuptake inhibitor</td>
</tr>
<tr>
<td>MAO inhibitor</td>
</tr>
<tr>
<td>Alpha-2 blocker</td>
</tr>
<tr>
<td>5-HT₁A activator</td>
</tr>
<tr>
<td>5-HT reuptake activator (tianeptine)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Adaptive Changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beta down-regulation</td>
</tr>
<tr>
<td>Alpha-1 up-regulation</td>
</tr>
<tr>
<td>5-HT₂ down-regulation</td>
</tr>
<tr>
<td>5-HT₁ sensitivity increase</td>
</tr>
<tr>
<td>D₂ sensitivity increase</td>
</tr>
<tr>
<td>D₁ down-regulation</td>
</tr>
</tbody>
</table>

NA: Norepinephrine
5-HT: Serotonin
MAO: Monoaminoxidase
D₂: Dopamine-2 receptors
D₁: Dopamine-1 receptors
It is not certain which group of antidepressants has the best effect on restoration of brain function. It is notable that substances with very different monaminergic efficacy have similar antidepressant properties. These include serotonin reuptake inhibitors, as well as serotonin reuptake blockers, which tend to lead to serotonin reduction rather than an increase [86]. Tianeptine, which lowers the serotonin concentration of receptors shows, for example, in addition to an antidepressive effect, definite reorganisational effects on stress-induced morphological changes in the hippocampus in animal models [87]. Thus tianeptine is possibly an interesting candidate for to promote brain recovery, but there are no related studies to date.

The TALOS-Trial in Denmark (so far published only as abstract) showed in a population of 642 stroke patients no significant effect of citalopram compared to placebo on brain recovery. There was only a slight statistical tendency for some functional improvement. Further large clinical studies (FOKUS AFFINITY EFFECT) are underway to clarify the effect of Fluoxetine on functional recovery 6 months after stroke in more than 6000 patients. Results are expected in 2018.

With respect to use of antidepressants, it can be stated that a higher-quality study following EBM criteria demonstrated that Fluoxetine has a definite effect on promoting restoration of brain function after a stroke.

Depression and Multiple Sclerosis

Many MS patients do not always immediately experience noticeable depression and should be treated with antidepressants after careful psychometric examination. The prevalence among depression among MS patients is between 47 and 54% [88], and the risk of suicide among these patients is twice as high compared to the normal population [89].

Impaired Consciousness

Various classes of substances have been used experimentally to improve the level of consciousness in patients who are unresponsive or exhibit minimal alertness. A positive effect has been shown in several small studies of amantadine [90–94].

A larger multi-centric randomized study demonstrated that amantadine was also effective following EBM criteria, even if only for a short period [71].

Somewhat positive effects were demonstrated for L-Dopa in smaller samples. Likewise, brain reorganization effects of other dopamine agonists such as pramipexole and bromocriptine were shown in small samples [94–97]. However, the dosage and duration of L-Dopa administration remains unclear.

Individual studies of pump-administered apomorphine indicated positive effects [98]. In a larger study of 80 patients, methylphenidate was shown to shorten the length of stay in intensive care as well as the entire length of hospitalization for those suffering from severe traumatic brain trauma [61]. However, Martin and Whyte in a study of 22 patients found that methylphenidate had no positive effects on improving consciousness [99]. Small studies have also shown positive effects for antidepressants such as sertaline [100] or classical tricyclic antidepressants such as amitriptyline [101]. Modafinil, a substance used predominantly for the treatment of parasomnia, has also shown some positive effects [102], but these were not confirmed by larger sample sizes [103].

Interestingly, the so-called ‘Z-drugs’ such as zopiclone and zolpidem, which are normally used for sleep stimulation and/or treating insomnia, show apparently paradoxical arousal effects. However such effects have only been observed in individual cases of coma treatment [104–109]. Due to the minimal potential for side effects, trial use of zolpidem for a few days may be useful for patients with unresponsive wakefulness as well as minimal conscious state (minimal responsive wakefulness).

In summary, apart from amantadine, a positive effect on the elevation of consciousness in states of restricted consciousness has not yet been demonstrated with sufficient certainty for any substance.

Fatigue

For MS patients fatigue is a particularly significant problem. Despite initial positive-appearing reports on the use of modafinil [110], its effects were not confirmed in further studies [111]. Likewise, other neurostimulants such as pemoline [112] and amantadine, as well as individual case observations on the potassium channel blocker fampridine, occasionally demonstrated effects [113–117]. Meta-analyses, however, could not confirm these results [118]. Taking alfalcacidol, an analog of vitamin D, may provide better effects [119]. Problems of treating fatigue in MS patients was recently extensively treated in a review [120].

Neuroprotective Substances

Despite good results in numerous animal experiments, the use of supposedly neuroprotective substances such as NMDA, AMPA receptor blockers, or calcium antagonists, etc., has been very disappointing. Despite hundreds of millions of dollars spent on in various studies, the results were generally completely negative [121–123].

Rogalewski et al. [123] had pointed out that the use of unimodal substances, that is, those acting on a 1:1 receptor-transmodal basis were less effective in achieving neuroprotective or neuroregenerative effects after stroke or cerebral trauma compared to pleiotropic multimodal substances.

Our recently published CARS trial using cerebrolysin was the first meaningful study following EBM criteria to demonstrate a verifiable clinically significant effect on stroke recovery when cerebrolysin was used during the subacute stage [124]. Among a total of 208 randomized patients, we were able to show that protracted administration of cerebrolysin (30 ml IV over 21 days) in combination with physiotherapy in the subacute phase (start of treatment 24-72 hours after onset) had a significant effect in an impairment measurement test (ARAT) even after three months post-stroke.

Cerebrolysin is a standardized mixture of low molecular weight neuropeptides and free amino acids and is thus a pleiotropic multimodal substance that can affect a variety of molecular processes of brain function repair.

Of course, this initial large study requires further substantiation by further investigations, especially when earlier studies, although smaller, showed only slight effects of the substance, e.g. [125, 126].
Cerebrolysin is certainly not a cure-all or even comparable to the legendary “heavenly drug” Therial of antiquity and Middle Ages which likewise consisted of a mixture of over 70 substances (see [127]).

To date cerebrolysin has not been approved as a medication in Germany.

Clinical Relevance

1. Selection of medication in patients in the subacute phase after stroke and cranial brain trauma should take into account basic aspects of the pharmacology of brain reorganization. In particular, so-called detrimental drugs should be avoided whenever possible and replaced by less harmful medications. This applies in particular to the use of certain anti-hypertensive and classical anticonvulsant substances such as phenytoin and barbiturates, but especially benzodiazepines and butyrophenones.

2. The use of antidepressants, in particular SSRIs, can also be recommended for the improvement of cerebral reorganization in non-depressed stroke patients as well.

3. The use of amantadine can be recommended to improve the level of consciousness in patients with impaired consciousness (non-responsive or minimal responsive alertness).

4. Of all the neuroprotective drugs, only the multimodal substance cerebrolysin has been shown to be clinically significant in the recovery of motor function in subacute stroke patients.

Conflict of Interest:

V. Hömberg has received honoraria from Ever Neuropharma Austria.

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