Does Dihydroergotamine Treat the "Whole Migraine"?

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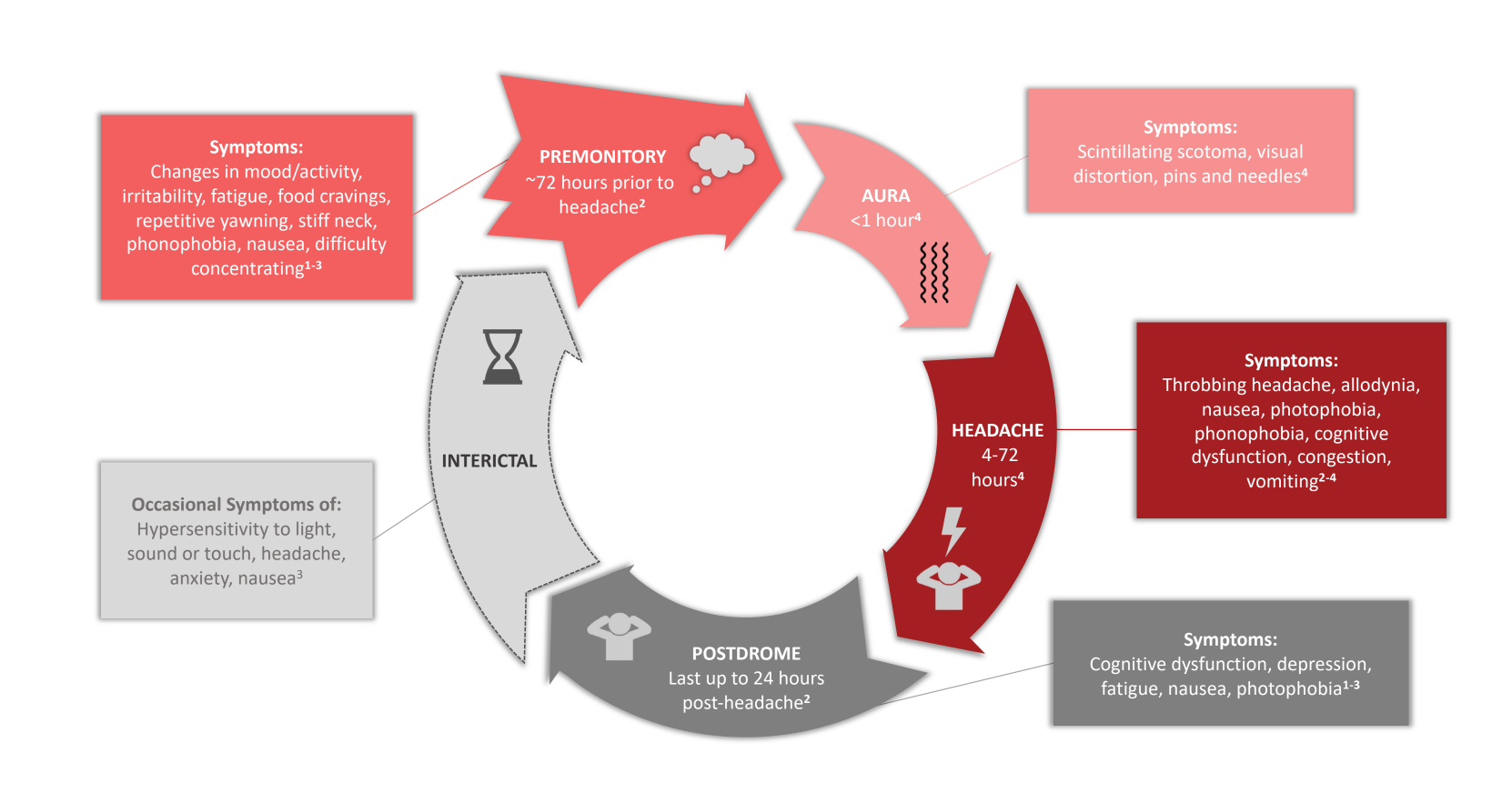
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Introduction

The Whole Migraine: Phases of the Migraine Cycle

 Migraine is a complex and multifaceted disorder with 5 distinct phases; therefore, it is important to consider treating the whole migraine (**Figure 1**)¹

Figure 1. The Five Phases of the Migraine Cycle Include the Premonitory, Aura, Headache, Postdrome, and Interictal Phases¹⁻⁴



Current Migraine Therapies and Limitations

- One of the primary goals for the treatment of migraine attacks includes the rapid relief of pain and associated symptoms⁵
- Most migraine therapies target a very narrow set of receptors focused mainly on headache pain
- Triptans are commonly used as acute medications and are 5-hydroxytryptamine (5-HT)_{18/1D} receptor agonists with some affinity for the 5-HT_{1F} receptor subtype⁶
- Novel emerging acute and preventive therapies include ditans (5-HT₁₅ receptor agonists), gepants (calcitonin gene-related peptide [CGRP] receptor antagonists), and anti-CGRP monoclonal
- Because migraine encompasses a spectrum of symptoms, this narrowly targeted receptor profile for most migraine therapies does not allow patients to achieve a holistic relief from migraine, and patients often discontinue treatment due to a lack of consistency, headache recurrence, and accompanying nausea^{1,5}

Dihydroergotamine (DHE)

- DHE has a long, established history as an effective migraine therapy and is well-regarded by physicians because of its^{7,8}:
- Rapid onset^{9,10}
- Efficacy against a full range of acute symptoms of migraine, including pain, photophobia, and phonophobia¹¹
- Efficacy irrespective of the time of treatment (**Table 1**)¹²

- DHE is effective in patients with difficult-to-treat migraine, such as those who⁸:
- Have status migrainosus⁷
- Wake with migraine¹³
- Are triptan resistant¹³
- Have menstrual migraine¹⁴
- Have allodynia^{11,13}
- Have severe or prolonged migraine^{9,11,13}
- There is also minimal risk of medication overuse with DHE use⁷

Table 1. The Pharmacokinetics of Different Routes of **Administration of DHE**

DHE Pharmacokinetics				
DHE Route of Administration	Dose (mg)	T _{max} (Minutes)	AUC _{0-2hours} (h*pg/mL)	Absolute Bioavailability*
Intravenous ^{7,15}	1	1-6	3019	100%
Intramuscular ^{7,16}	1	15-24	4791	100%
Subcutaneuous ^{8,17}	1	20-40	N/A	100%
Nasal spray ⁷	2	48-60	428.7-1316	32%†
POD ^{®7,18}	1.45	30	1595	59%

AUC, area under the curve; DHE, dihydroergotamine; N/A, not available; POD, precision olfactory delivery. *Absolute bioavailability is defined as the amount of drug from a formulation that reaches systemic circulation relative

†Nasal spray refers to Migranal®, which has a variable bioavailability; 32% according to the prescribing information and 13-101% according to the Summary Basis of Approval. 19

Methods

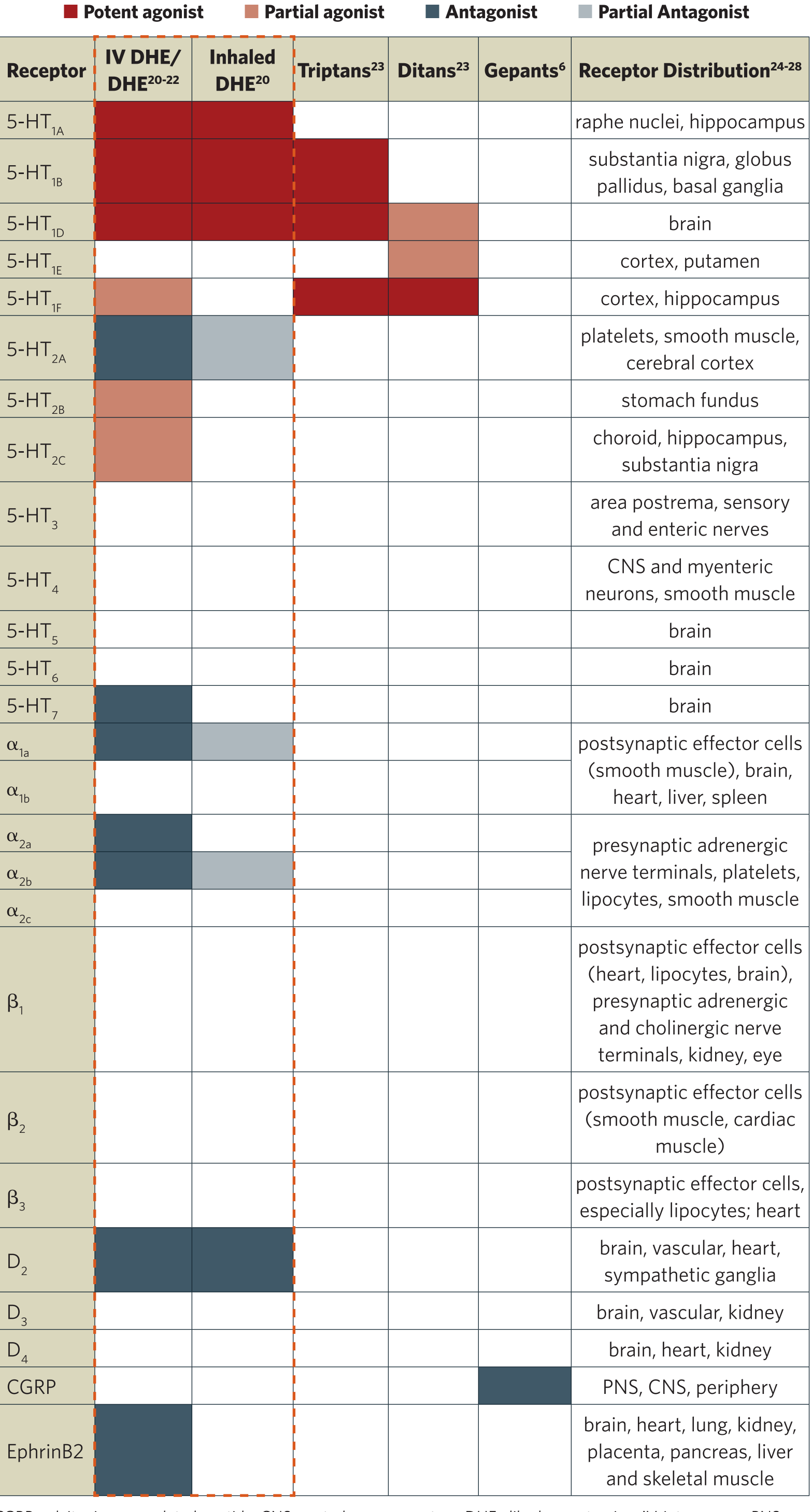
This is a review of the comparative pharmacology of acute treatments for migraine performed by conducting a literature review of the pharmacology and biological activity of new and existing migrainespecific treatments

Objectives

- To compare receptor pharmacology of acute treatments for migraine in published literature
- To examine the pharmacology of acute treatments for migraine in the context of the migraine cycle to provide insight into the total migraine benefits of treating with a broad receptor binding agent, such as DHE, versus the limitations of treating with narrowly targeted therapies
- To find a possible explanation for why DHE has such a high response rate even in the most difficult-to-treat migraine

Results

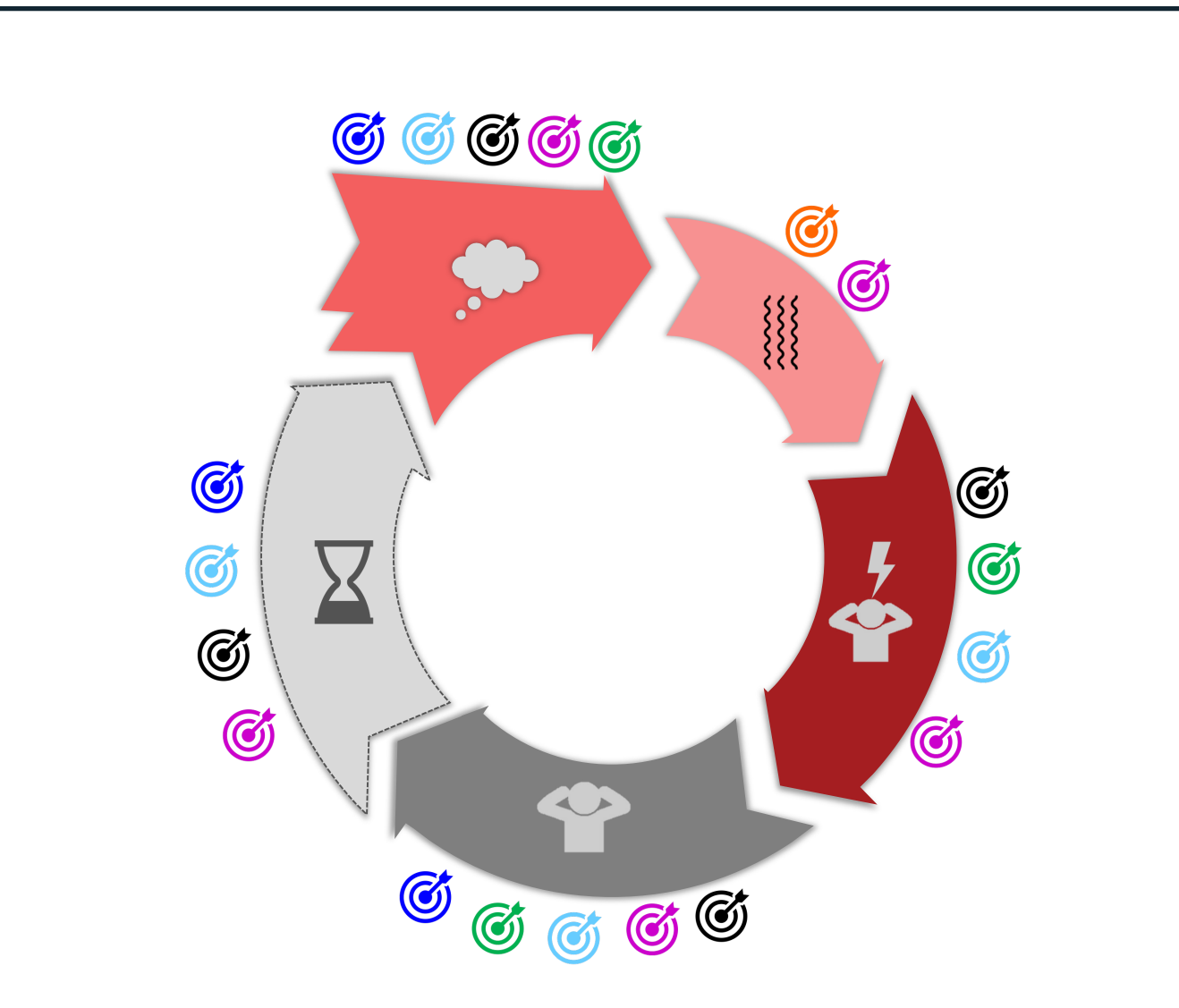
Table 2. Compared to Other Acute Migraine Therapies, DHE Displays a Broad Range of Pharmacological Activity at Serotonergic (5-HT), Adrenergic (α), and Dopaminergic (D) Receptor Subtypes



CGRP, calcitonin gene-related peptide; CNS, central nervous system; DHE, dihydroergotamine; IV, intravenous; PNS, peripheral nervous system. Note: Functional receptor binding data presented in this table is not a direct comparison, but a review of available,

published data on functional receptor binding for acute migraine drug therapies. Inhaled DHE meslyate is MAPOOO4

Figure 2. Hypothetical Model for How DHE Targets the Whole Migraine



- The can alleviate headache pain symptoms via several mechanisms
 - Constricts pain-producing intracranial extracerebral blood vessels via 5-HT_{1R} receptors¹⁶
 - Inhibits trigeminal neurotransmission via activation of peripheral and central 5-HT_{1D} receptors¹⁶
 - Effective in patients with allodynia¹¹
 - Agonist of 5-HT₁, receptors, which may have a beneficial role in migraine prophylaxis, which may lengthen the interictal period²⁰
 - Central effects of DHE may reverse central sensitization⁸
 - Antagonist of 5-HT $_{2\Delta}$ receptors, a pronociceptive receptor that can increase headache attacks²⁹
 - It has been suggested that antimigraine effects of DHE involve repressing CGRP release via activation of $\alpha 2$ -adrenoceptors³⁰
- May confer anti-nausea activity via D₂ receptor antagonism²⁰
- Is effective for acute treatment of migraine with aura 12
- (♂) Has 5-HT₂₄ receptor antagonism, which is involved in mood disorders³⁰
- Exerts D₂ antagonism, and a dopamine imbalance during migraine attacks may contribute to pain, discomfort, increased sensory sensitivity, and aversive reactions to environmental stimuli³²
- Binds to the ephrinB2 receptor which has been implicated in memory, and DHE has been shown to improve memory problems in post-concussive syndrome^{33,34}

Conclusion

- Unlike other migraine therapeutics, DHE interacts with several receptor families and subtypes, which include serotonergic, adrenergic, and dopaminergic subtypes (Table 2)
- DHE also slowly dissociates from 5-HT_{1R/1D} receptor sites, which may explain why DHE has sustained anti-migraine effects³⁵
- DHE is able to exert a greater influence than single receptor agonists/ antagonists over the pathophysiology of the migraine cycle due to its widespread pharmacological activity (Figure 2)
- DHE administered at consistent doses and optimal plasma concentrations not only maximizes therapeutic gain, but also improves safety and tolerability^{7,20}
- Advances in DHE delivery systems will further address these issues^{7,20}

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Disclosures and Acknowledgments

All authors are full-time employees and stockholders of Impel NeuroPharma. This research was sponsored by Impel NeuroPharma. Editorial support was provided by IMPRINT Science, and funded by Impel NeuroPharma.

