The identification of noninvasive biomarkers for diagnosis and prognosis is a fundamental challenge in current schizophrenia (ScZ) research (1). Importantly, biomarkers should ideally allow insights into the underlying pathophysiological mechanisms and facilitate links to preclinical research (2). One potential candidate is the 40-Hz auditory steady-state response (ASSR). ASSRs reflect stimulus rate–dependent, evoked activity to constant periodic stimuli in different sensory modalities that can be detected using electroencephalography (EEG) and magnetoencephalography (MEG). ASSRs show a peak frequency around 40-Hz in humans (3) in contrast to other sensory modalities, such as visual SSRs (4).

Kwon et al. (5) investigated 40-Hz ASSRs for the first time in patients with ScZ by using EEG, demonstrating reduced power and phase delay to 40-Hz stimulation. These findings provided further support for the hypothesis that neural circuits were compromised in ScZ to generate oscillations in the gamma band (30–100 Hz) (6). During normal brain functioning, gamma-band oscillations have been proposed to facilitate coordination of distributed neuronal activity to support the generation of perception and cognition (7) and are accordingly a candidate mechanism for the pervasive sensory and cognitive deficits in ScZ (8).

The initial findings by Kwon et al. (5) have been replicated by several groups using both EEG and MEG (9–11). In addition, more recent studies have examined 40-Hz ASSRs in participants at clinical high risk for psychosis (CHR-P) and first-episode psychosis (FEP) patients (12,13), raising the possibility that 40-Hz ASSRs could be used for early detection and diagnosis of early-stage psychosis.

Despite the extensive evidence of 40-Hz ASSR deficits in ScZ, several questions remain regarding their significance and interpretation. Therefore, the goal of this article is to provide a state-of-the-art overview of 40-Hz ASSRs as a mechanistic biomarker for elucidating circuit dysfunctions in ScZ as well as for early detection and diagnosis. Accordingly, we will summarize the mechanisms underlying 40-Hz ASSRs, drawing on computational, physiological, and pharmacological perspectives. This will be followed by an overview of current studies in CHR-P participants, FEP groups, and patients with ScZ. Finally, evidence from genetics as well as recent studies on 22q11.2 deletion syndrome will be reviewed together with data on developmental modifications, followed by recommendations for future research.
40-Hz ASSR, in particular the medial Heschl’s gyrus (14,15), more recent work has suggested that 40-Hz ASSRs involve a more extensive network. Evidence for the contribution of frontal generators toward 40-Hz ASSRs in humans comes from MEG/EEG (16,17) as well as from intracranial recordings (18) (Figure 1). In addition, 40-Hz ASSRs have been observed in parietal areas (14) and in the inferior colliculus (19). Tada et al. (18) analyzed high-density electrocorticography in response to ASSRs at 20, 30, 40, 60, 80, 120, and 160 Hz using two common techniques to analyze steady-state activity, intertrial phase coherence (ITPC) (20) and spectral power estimates. The first refers to the consistency of phase angles across trials, therefore reflecting only evoked activity. The latter encompasses both evoked activity and induced components, of which the timing can differ between trials. Modulation of ITPC and spectral power was maximal at 40-Hz stimulation and was distributed across temporal, parietal, and frontal cortices.

In addition, there is evidence for a role of the thalamus, in particular the medial geniculate body, in the generation of 40-Hz ASSRs from MEG/EEG, positron emission tomography, and functional magnetic resonance imaging data (12,19,21). These findings were corroborated by a study showing that electrical stimulation of thalamic neurons evoked gamma-band activity around 40-Hz in the auditory cortex (22). Moreover, recent work has shown that generators extend to additional subcortical areas, including the hippocampus (12) and brainstem (15).

CIRCUIT MECHANISMS OF 40-HZ ASSRs: EXCITATION/INHIBITION BALANCE PARAMETERS

Unlike transient evoked potentials, SSRs require a high temporal resolution for coordinated signal integration as well as for transmission and processing, especially in higher-frequency ranges (23). Initial efforts focused on disclosing whether SSRs simply constitute a summation of event-related potentials or whether they reflect the entrainment of rhythmic oscillatory activity. While some studies supported the summation hypothesis (24,25), it is more likely that both models are nonexclusive and interacting to produce the ASSR response.

As such, it has been proposed that early transient components of the ASSR response may reflect event-related potential processes, while late-latency, sustained responses reflect rhythmic activity (11,16). This theory is supported by evidence indicating that ITPC is larger for the late sustained 40-Hz ASSRs (150–500 ms) compared with ITPC modulation between 0 and 50 ms (12,26), suggesting that sustained rhythmic activity may be only observed after the early evoked component. In this context, it is important to highlight that neural oscillations reflect synchronous, rhythmic activity of neuronal ensembles that occur in a circumscribed frequency range and is sustained over several cycles (27). Accordingly, neural oscillations need to be distinguished from broadband power changes, transient responses, and aperiodic activity (28), and recent methods have been introduced to separate these processes (29).

Gamma-band oscillations emerge from the balance between excitation and inhibition (E/I) in neural networks (30) (Figure 2). Specifically, the time constants of inhibitory post-synaptic potentials of GABAergic (gamma-aminobutyric acid-ergic) parvalbumin-positive (PV+) interneurons are ideally suited to generate 40-Hz rhythms (31). This has been shown, for instance, through hippocampal excitation of PV+ interneurons by means of NMDA receptors (NMDARs), which resulted in 40-Hz transient oscillatory responses in pyramidal cells (32,33).

PV+ interneurons mainly target the perisomatic region of pyramidal cells and can therefore control their output effectively as opposed to somatostatin-expressing interneurons that mainly inhibit the apical dendrite (34). Computational studies have further demonstrated that these anatomical and electrophysiological properties of PV+ interneurons are crucial for the generation of gamma-band oscillations and that several key E/I balance parameters determine their power and coherence (35). Moreover, the strength of exerted inhibition, which in turn is dependent on the strength of NMDAR activation as well

Figure 1. 40-Hz auditory steady-state responses (ASSRs) in electroencephalographic/magnetoencephalographic (EEG/MEG) data. (A) 40-Hz auditory stimulation elicits SSRs measurable with EEG/MEG. Illustrated are typical response patterns in both 40-Hz power (analyzed with time–frequency analysis [TFA]) and phase (analyzed with intertrial phase coherence [ITPC]). In EEG recordings, these responses can be observed over frontocentral regions, while in MEG they are localized over temporal regions. (B) 40-Hz ASSRs consist of early transient and late sustained activity; illustrated are a typical neuronal ASSR of human recordings using MEG, depicted by ITPC values and the averaged steady-state potential. The difference between the early onset and the late sustained activity are clearly visible. (C) Overview of generators involved in 40-Hz ASSRs from multimodal imaging studies using positron emission tomography/functional magnetic resonance imaging/EEG/MEG and intracranial recordings. (From P.J. Uhlhaas, Ph.D., et al., unpublished data, November 2022.)
as on the maximal conductance of the GABAergic synapses, also crucially influences 40-Hz ASSR power (36–39).

Further evidence for the role of E/I balance parameters in the generation of 40-Hz ASSRs comes from pharmacological studies. In human EEG recordings, the NMDAR antagonist ketamine has been associated with an increase in the power of the 40-Hz ASSRs (40). In contrast, administration of the NMDAR antagonist MK 801 into the medial geniculate body in mice was associated with reduced 40-Hz ASSRs in the auditory cortex, without affecting the early transient response (41). Such diverging findings could be explained by different locations and dosages of drug administration. For instance, Sivarao et al. (42) showed that NMDAR channel occupancy is related to the modulation of both power and phase locking of 40-Hz ASSRs, with lower ketamine dosages causing an increase in spectral power and ITPC, while higher doses caused decreased 40-Hz ASSRs.

Similarly, there is emerging evidence that GABAergic neurotransmission modulates 40-Hz ASSRs. Increasing inhibition via administration of the GABAA agonist muscimol results in increased power and phase locking of 40-Hz ASSRs in humans (43). Moreover, selective PV+ interneuron excitation using optogenetic stimulation in the basal forebrain in rats increased ASSR responses in the auditory cortex only when stimulating at 40 Hz but not at other frequencies (44). Accordingly, these data indicate that modulation of both power and ITPC values of 40-Hz ASSRs are a sensitive marker for E/I balance alterations that could allow the identification of circuit mechanisms in ScZ. In particular, both PV+ interneuron activation and the excitatory drive mediated through NMDARs have a mechanistic impact on 40-Hz ASSRs.
40-HZ ASSRs IN ScZ: PATTERN OF DEFICITS AND CORRELATIONS WITH CLINICAL VARIABLES

Approximately 40 studies have investigated 40-Hz ASSRs in patients with ScZ using both EEG and MEG [for a recent review, see (44)]. The large majority of studies have reported a reduction in both ITPC and power of 40-Hz ASSRs with medium-level effect sizes ([6], but see [45,46]). This pattern is consistent with evidence from other sensory and cognitive paradigms in ScZ ([6]), suggesting that neural circuits involved in the generation of high-frequency oscillations are impaired.

So far, however, studies have focused almost exclusively on the analysis of sensor-level data and source localization of auditory regions as the origin of 40-Hz ASSRs deficits. Accordingly, it is unclear which areas are fundamentally implicated beyond the auditory cortex given the contribution of extensive cortical and subcortical regions toward the generation of 40-Hz ASSRs ([12,18]). Koshiyama et al. (47) applied a Granger causality analysis, a functional connectivity measure, to assess the propagation of 40-Hz ASSRs across cortical sources in a large sample of patients with ScZ and control participants. Patients with ScZ showed a complex pattern of increased and decreased connectivity across the early transient as well as during the later sustained responses that involved temporal and frontal brain regions.

In patients with ScZ, there is evidence for a reduction in both the early transient and sustained 40-Hz ASSRs (48) that may, however, differ across early versus later illness stages (49). In addition, Kwon et al. (5) reported a delay between click onset and the subsequent negative peak in band-filtered time domain EEG data. This finding was replicated by Roach et al. (60) [see also (51)], highlighting that the phase delay deficit in patients with ScZ was associated with a significantly larger effect size than both spectral power and ITPC reductions.

An important aspect concerns the specificity of ASSR deficits toward 40-Hz stimulation. While auditory cortices during normal brain functioning respond preferentially to 40-Hz ASSRs (18), there is consistent evidence that impairments in ScZ extend to other frequencies. Thus, ASSR deficits have also been observed at 80 Hz but not at 20 or 30 Hz (52). In addition, several studies have shown that ASSRs at both delta (1–4 Hz) (53) and theta (4–7 Hz) (54) bands are reduced in ScZ as well. During normal brain functioning, there is evidence that low- and high-frequency oscillations interact: for example, the amplitude of gamma-band activity can be modulated by the phase of low-frequency (delta/theta-band) oscillations (55). However, studies that investigated cross-frequency coupling showed that 40-Hz ASSR deficits were not related to lower frequencies in ScZ (56,57).

Among the clinical correlates, correlations between increased 40-Hz ASSRs and elevated positive symptoms, especially auditory hallucinations (58,59), have been reported, which, however, has not been confirmed by other studies (52). In addition, Oguy et al. (60) examined whether 40-Hz ASSRs differentiated patients with ScZ who did not respond to standard antipsychotics (treatment-resistant schizophrenia [TRS]) versus a non-TRS group. Evoked power during 40-Hz ASSRs was only impaired in the TRS group compared with the control group. However, no differences were found between TRS and non-TRS patients in 40-Hz ASSR power.

Given that gamma-band oscillations have been proposed to underlie impaired cognitive and sensory processes in ScZ ([8]), correlations between deficits in 40-Hz ASSRs, cognition, and possibly also functional impairments can be expected. Robust relationships with cognitive deficits have not been demonstrated so far ([12,56,61]). With regard to functional impairments, there is preliminary evidence that reduced 40-Hz ASSRs correlate with lower functional status in patients with ScZ (61).

Finally, several studies have examined the relationship between anatomical alterations, especially gray matter volume, and 40-Hz ASSRs in ScZ. Thus, there is evidence that gray matter reductions in the auditory cortex correlate with decreased 40-Hz ASSRs (62). A study by Du et al. (63) that linked resting-state functional magnetic resonance imaging data to sensor-level 40-Hz ASSRs suggested, however, that a network of brain areas consisting of the temporal medial prefrontal cortex and postcentral/precentral gyrus is associated with deficient 40-Hz ASSRs.

40-HZ ASSR DEFICITS IN EARLY-STAGE PSYCHOSIS

More recent work has investigated whether 40-Hz ASSR impairments are present during early-stage psychosis to address the potential as a biomarker for early detection and diagnosis (Table 1). This is particularly important, as early intervention can modify the trajectory of patients with FEP (64), and there is an urgent need for biomarkers to stratify patients according to clinical outcomes and pathophysiological mechanisms (65).

Currently, several studies have investigated 40-Hz ASSRs in patients with FEP (12,13,49,58,66–68), the majority of which reported robust impairments in both spectral power and ITPC, while Bartolomeo et al. (66) and Coffman (58) found intact 40-Hz power. With regard to CHR-P participants, Lepock et al. (69) showed intact ITPC and 40-Hz ASSR power, while Grent’-t-Jong et al. (12), Koshiyama et al. (66), and Tada et al. (49) found evidence for impaired 40-Hz ASSRs.

Furthermore, Grent’-t-Jong et al. (12) examined the question of whether 40-Hz ASSRs could constitute a biomarker for clinical outcomes in CHR-P participants, such as persistence of attenuated psychotic symptoms and transition to psychosis (Figure 3). Source-reconstructed 40-Hz ASSRs revealed that both the CHR-P and FEP groups had an overlapping deficit in spectral power and ITPC in the auditory cortex, hippocampus, and thalamus. Importantly, both attenuated psychotic symptom persistence and transition to psychosis were predicted by 40-Hz ASSR deficits in the hippocampus, thalamus, and superior temporal gyrus.

Several studies also examined associations between 40-Hz ASSRs, functioning, and symptoms in early-stage psychosis. Similar to findings in established ScZ, however, correlations with functioning (12,70), symptoms (12,58), and cognitive impairments (12) were inconsistent across studies.

40-HZ ASSRs, GENETICS, AND BRAIN DEVELOPMENT

The heritability of ScZ is estimated at approximately 80% (71), and recent genome-wide association studies have identified risk genes that impact on E/I balance parameters (72). There is
Auditory 40-Hz Steady-State Responses in Schizophrenia

Table 1. 40-Hz ASSR Studies in Participants With FEP and Participants at CHR for Psychosis

<table>
<thead>
<tr>
<th>Study</th>
<th>Clinical Group</th>
<th>Age, Mean (SD)</th>
<th>Recording Technique</th>
<th>Stimulus Type</th>
<th>Main Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ahmed et al. (70)</td>
<td>35 CHR-P</td>
<td>21 (3.4)</td>
<td>32-channel EEG</td>
<td>Click trains 40 Hz</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>34 HC</td>
<td>28 (8.7)</td>
<td>60-channel EEG</td>
<td>Click trains 20, 30, and 40 Hz 500-ms duration</td>
<td>FEP-SZ &lt; HC: power and ITPC</td>
</tr>
<tr>
<td></td>
<td>16 FE-SZ</td>
<td>26 (8.1)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>16 FE-AF</td>
<td>24 (7.4)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tada et al. (49)</td>
<td>21 HC</td>
<td>22 (3.3)</td>
<td>64-channel EEG</td>
<td>Click trains 20, 30, and 40 Hz 500-ms duration</td>
<td>CHR-P &lt; HC: only late-latency power and ITPC</td>
</tr>
<tr>
<td></td>
<td>15 CHR-P</td>
<td>22 (4.0)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>13 FEP</td>
<td>25 (5.9)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Koshiyama et al. (86)</td>
<td>24 HC</td>
<td>22 (3.0)</td>
<td>64-channel EEG</td>
<td>Click trains 20, 30, and 40 Hz 500-ms duration</td>
<td>CHR-P &lt; HC: only late-latency power and ITPC</td>
</tr>
<tr>
<td></td>
<td>27 CHR-P</td>
<td>21 (3.9)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>21 ROSZ</td>
<td>24 (6.7)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wang et al. (67)</td>
<td>28 HC</td>
<td>26 (5.5)</td>
<td>64-channel EEG</td>
<td>Click trains 40 Hz 500-ms duration</td>
<td>FEP &lt; HC: power and ITPC</td>
</tr>
<tr>
<td></td>
<td>33 FEP</td>
<td>25 (6.6)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bartolomeo et al. (68)</td>
<td>19 HC</td>
<td>22 (4.3)</td>
<td>28-channel EEG</td>
<td>Click trains 40 Hz 500-ms duration</td>
<td>FEP = HC: power</td>
</tr>
<tr>
<td></td>
<td>34 FEP</td>
<td>23 (3.6)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lepock et al. (69)</td>
<td>22 HC</td>
<td>22 (3.0)</td>
<td>32-channel EEG</td>
<td>Click trains 40 Hz 500-ms duration</td>
<td>CHR-P = HC: power and PLF</td>
</tr>
<tr>
<td></td>
<td>36 CHR-P</td>
<td>21 (3.4)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grent-Jong et al. (12)</td>
<td>49 HC</td>
<td>23 (3.6)</td>
<td>248-channel MEG</td>
<td>AM sounds 40 Hz 2000-ms duration</td>
<td>CHR-P &lt; HC: ITPC in RHES, power in RTHA and RHIP</td>
</tr>
<tr>
<td></td>
<td>38 CHR-N</td>
<td>23 (4.7)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>116 CHR-P</td>
<td>22 (4.5)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>33 FEP</td>
<td>24 (4.5)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coffman et al. (58)</td>
<td>32 HC</td>
<td>24 (5.5)</td>
<td>63-channel EEG</td>
<td>Click trains 40 Hz 500-ms duration</td>
<td>FEP = HC: power and ITPC</td>
</tr>
<tr>
<td></td>
<td>25 FEP</td>
<td>24 (4.0)</td>
<td></td>
<td></td>
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</tbody>
</table>

AM, amplitude modulated; ASSR, auditory steady-state response; CHR-P, clinical high risk for psychosis; CHR-N, negative clinical high risk; EEG, electroencephalography; FE, first-episode psychosis; FE-SZ, first-episode schizophrenia; HC, healthy control; ITPC, intertrial phase coherence; MEG, magnetoencephalography; PLF, phase-locking factor; RHES, right Heschl’s gyrus; RTHA, right hippocampus; ROSZ, recent-onset schizophrenia; RTHA, right thalamus; UHR, ultra high risk.

*If more stimulation frequencies were presented, only the results from the 40-Hz stimulation condition are reported here.

preliminary evidence that unaffected first-degree relatives exhibit reductions in 40-Hz ASSRs (73). Moreover, computational modeling has shown the impact of common variants on 40-Hz ASSRs (74), suggesting that 40-Hz ASSR deficits are closely linked to genetic risk and therefore could constitute an endophenotype. This possibility is consistent with data showing that 40-Hz ASSR deficits can also be found in conditions that are characterized by overlapping circuit dysfunctions and genetics, such as bipolar disorder and autism spectrum disorder (75–78).

More recent studies have examined the relationship between 40-Hz ASSR deficits in 22q11.2 deletion syndrome (17,79), which is a neurogenetic disorder that confers a 30% to 40% lifetime risk for the development of psychosis (80). Several genes within the 22q11.2 region have been linked to glutamatergic and GABAergic neurotransmission (81) and disrupted migration and placement of cortical interneurons (82). Mancini et al. (17) examined 40-Hz ASSRs in EEG data in a sample of 22q11.2 deletion carriers and control participants (Figure 4). Both the power and ITPC of 40-Hz ASSRs as well as the evoked theta-band power were impaired in deletion carriers. Gamma-band spectral power reductions were particularly prominent in the anterior cingulate cortex, posterior cingulate cortex, thalamus, and right primary auditory cortex. Moreover, 40-Hz ASSR deficits were pronounced in deletion carriers with psychotic symptoms and correlated with the reduction of gray matter in the auditory cortex. Importantly, a linear increase of 40-Hz ASSR spectral power was observed from childhood to adulthood in healthy control participants but not in deletion carriers.

There is converging evidence that E/I balance, such as PV interneurons and their excitatory inputs (83), as well as GABAergic subunits (84), undergo important modifications during adolescence, which could in turn provide not only sensitive periods for risk factors, such as cannabis (85), but also interventions that could potentially modify and even restore existing circuit dysfunctions. A study by Mukherjee et al. (86) examined the possibility to modify PV+ interneuron functioning in a mouse model of 22q11.2 deletion syndrome during development. Adult mice were characterized by low PV+ interneuron plasticity as well as by pronounced deficits in cognitive tasks and gamma-band oscillations. Importantly, cognitive dysfunction could be prevented permanently by dopaminergic D2 receptor antagonist treatment or by chemogenetic activation of PV+ interneurons during late adolescence.
40-HZ ASSR DEFICITS IN ScZ: LINK TO E/I BALANCE CIRCUIT DYSFUNCTIONS

Impaired gamma-band oscillations in ScZ have been linked to altered E/I balance parameters, in particular PV+ interneuron deficits (87) as well as NMDAR hypofunctioning (88). However, the precise pattern and contributions of NMDARs and GABAergic interneurons toward impaired gamma-band oscillations in ScZ remain unclear. One possibility is that circuit deficits are due to a primary dysfunction in inhibitory interneurons in ScZ (87). In addition, evidence exists that impaired inhibition could be the result of NMDAR hypofunctioning on PV+ interneurons (89) or reduced NMDAR drive on pyramidal cells (90).

A shift toward increased excitation as a result of NMDAR hypofunctioning has been implicated in circuit dysfunctions in ScZ, in particular during early-stage psychosis (91). Ketamine, an NMDAR antagonist, is associated with disinhibition in local (92) and large-scale networks (90) and with a dysregulation of gamma-band oscillations (93). Specifically, lower dosages of NMDAR antagonists, which elicit psychomimetic effects in healthy volunteers, cause preferentially an upregulation of both NMDAR antagonists, which elicit psychomimetic effects in healthy volunteers, cause preferentially an upregulation of both NMDAR antagonist, is associated with disinhibition in local (92), CHR-P (blue), and FEP (red) groups. (C) Regions of interest for which virtual channel data were computed and statistically examined for group differences. (D) Cumming estimation plots with data distribution swarm plots and group difference data, including HC participants and patients with FEP and three subgroups of CHR-P participants, based on 1-year follow-up data: nonpersistent (remitted) attenuated psychotic symptoms (APS-NP), persistent APS (APS-P), CHR-P participants who transitioned during follow-up time (CHR-P-T) (up to 36 months). Linear discriminant analysis predicts clinical outcomes of CHR-P individuals based on magnetencephalography data for APS-P vs. APS-NP: for the right hippocampus (RHIP), right middle temporal gyrus, and right superior temporal gyrus (RSTG) ITPC, area under the curve = 0.845. A cross-validated total of 27 (89.2%) of 39 APS-NP and 25 (73.5%) of 34 APS-P participants were correctly classified. Classification of CHR-P-T vs. nontransitioned CHR-P participants: for the right thalamus (RTHA), area under the curve = 0.695. Cross-validated totals of 52 (53.6%) of 97 in the nontransitioned CHR-P group and 10 (76.9%) of 13 in the CHR-P-T group were correctly classified. These plots show the predictive value of 40-Hz auditory steady-state responses in CHR-P individuals. [Adapted with permission from Grent-’t-Jong et al. (12).]
with recent postmortem data (100) that have indicated that reductions in PV+ interneurons may constitute an adaptive response to decreased excitatory drive.

Thus, the current findings suggest that 40-Hz ASSR deficits in ScZ may be compatible with altered E/I balance parameters. Specifically, evidence from pharmacology, postmortem studies, and computational modeling converges on the notion that reduced 40-Hz ASSRs in ScZ could be the result of impaired PV+ interneuron functioning (44,97). However, it is currently unclear whether this dysfunction is primary or secondary to an excitatory deficit.

While the concept of E/I imbalance has been useful to gain a first mechanistic understanding of circuit deficits in ScZ, current models of E/I balance should be extended to capture important aspects of the ASSR response and generation. For example, most models do not capture the diversity of cortical interneurons, with some exceptions (97,101). Furthermore, modeling studies so far have assumed that the ASSR response can be considered a simple summation of event-related potentials and have not addressed the entrainment of ongoing intrinsic oscillations by periodic ASSR stimuli.

**RECOMMENDATIONS FOR FUTURE RESEARCH**

First, it has been found that 40-Hz ASSRs are influenced by a number of experimental parameters that require careful consideration. Attention has been shown to modulate 40-Hz ASSR power in healthy control participants (102) but not in patients with FEP (58) or in patients with ScZ (103). Accordingly, future studies need to examine and control more carefully differences in attention as a possible confound for 40-Hz ASSR deficits in ScZ.

Second, state-dependent variables, such as arousal, may impact the strength of 40-Hz ASSRs (104). This is supported by findings indicating that eyes-open versus eyes-closed conditions modulate the strength of 40-Hz ASSRs (105). It is currently unclear, however, whether these manipulations differ in ScZ or FEP groups (67,106). Finally, it has been argued that amplitude-modulated sounds, compared with click trains, are more powerful in detecting late sustained 40-Hz impairments, whereas click trains are most sensitive to detecting early-lateness deficits (107). However, studies using click-train paradigms have shown sustained impairments for the duration of stimulation (9).

Regarding EEG/MEG analytic approaches, the consideration of baseline differences deserves careful consideration. Kim et al. (46) showed that 40-Hz ASSR impairments in patients with ScZ could only be found when the data were not baseline normalized, suggesting that higher noise levels may be present. In addition, the involvement of brain regions beyond the auditory cortex, such as the thalamus, hippocampus, and frontal regions, is not reflected in the large majority of current EEG/MEG studies. Given the extensive contribution of extended cortical and subcortical networks...
toward the generation of 40-Hz ASSRs \((12,18,108)\), future analytic protocols should ideally apply whole-brain source-localization approaches. A recent study by Grent'-'t-Jong et al. \((12)\), for example, showed that specifically subcortical generators in the thalamus and hippocampus were strongly impaired in CHR-P and FEP groups and that reduced 40-Hz ASSRs in the thalamus predicted transition to psychosis in CHR-P participants.

Finally, given that 40-Hz ASSRs reflect E/I balance parameters and that deficits in ITPC and spectral power may constitute a transdiagnostic biomarker, future studies could examine 40-Hz ASSRs across ScZ, bipolar disorder, and autism spectrum disorder to define potentially novel illness subtypes \((109)\) involving E/I balance parameters.

**SUMMARY**

The available evidence suggests that 40-Hz ASSRs are a promising biomarker, which is robustly impaired in patients with ScZ across different paradigms and recording modalities \((9,10)\). In addition, test-retest reliability has been established in several studies \((110,111)\). Importantly, there is also emerging evidence that both CHR-P and FEP groups are characterized by similar impairments in ITPC and spectral power during 40-Hz stimulation \((12,49)\) that could be potentially relevant for early detection and diagnosis. Developmental data furthermore indicate that 40-Hz ASSRs and the underlying generating mechanisms undergo major modifications during adolescence \((17,83,84)\), indicating a sensitive period not only for risk factors, but also for intervention to correct circuit anomalies.

Moreover, the 40-Hz ASSR also fulfills the criteria for a translational and mechanistic biomarker. Data from animal work have shown that similar to findings obtained from EEG and MEG recordings in humans, 40-Hz ASSRs elicit similar perturbations in both spectral power and ITPC that can be linked to circuit mechanisms fundamentally implicated in ScZ, in particular GABAergic interneurons and NMDARs. Together with computational modeling \((87,95)\), these data allow the testing of mechanistic hypotheses that could lead to the development of targeted and more effective interventions.

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Auditory 40-Hz Steady-State Responses in Schizophrenia


