ABSTRACT

Background: The last decade has seen development of numerous novel antipsychotic drugs with unique mechanisms including long-acting formulations for clinical use. A comparative assessment of these new drugs with each other and previous antipsychotics have not been performed with regards to risk for drug-induced movement disorders (DIMD).

Methods: Medline was searched from January 2010 to February 2022 for primary research articles and review articles in English using the search terms “extrapyramidal” and “tardive” with individual drug names of novel antipsychotics.

Results: We identified articles describing the risk of DIMD with 6 novel antipsychotics, 4 novel formulations, and 3 experimental antipsychotics. Both short- and long-term data generally showed comparable to lower risk of DIMD with novel antipsychotics and recent long-acting formulations compared to previously marketed antipsychotics.

Discussion: Several novel antipsychotics, particularly lumateperone and pimavanserin, show promise in being able to treat psychosis while reducing the risk of DIMD. Long-acting paliperidone may reduce risk of DIMD while other long-acting injectable formulations of SGA have similar risk of DIMD compared to oral formulations. New drug targets for treating psychosis without dopamine blockade also show promise.
INTRODUCTION TO DRUG-INDUCED MOVEMENT DISORDERS (DIMD)

The association of antipsychotics with DIMD has been attributed primarily to their dopamine 2 receptor (D2R) blocking properties [1]. Acute DIMD refers to early-onset and reversible phenomena including akathisia, dystonia, parkinsonism, and tremor [2]. Tardive DIMD have a variable presentation, most commonly including stereotypy or chorea, but also dystonia, akathisia, myoclonus, tics, and tremor occurring at least weeks to months after drug exposure and often persisting despite drug cessation [1, 3]. The prevalence of tardive syndromes among antipsychotic-treated patients is estimated to be between 20–30%, with annual incidence significantly increased by exposure to first-generation antipsychotics (FGAs) versus second-generation antipsychotics (SGAs) of 5.4–7.7% versus 0.8–3% (p < 0.0001), respectively, as well as by patient age [4, 5]. Although SGA use has reduced the incidence of DIMD compared to FGAs, the expansive marketing and prescription of SGAs may increase the absolute number of individuals at risk [6]. Understanding how novel antipsychotics influence the risk of the development of acute and tardive DIMD is critical to reducing patient harm. Although comparisons between reported studies are constrained by differences in design, methodology, and sample populations, we herein descriptively review data on acute and tardive DIMD associated with novel antipsychotics that have received Food and Drug Administration (FDA) approval since 2010 in addition to drugs in development.

METHODS

In this review, we restricted medications discussed to antipsychotic agents approved by the FDA since 2010: brexpiprazole, cariprazine, lumateperone, lurasidone, paliperidol, aripiprazole, pimavanserin, olanzapine-samidorphan (O-S), long-acting injectable (LAI) risperidone, LAI paliperidone, transdermal asenapine, and inhaled loxapine. The terms in the Medline database searched from January 2010 to February 2022 and the identified additional articles through reference review are indicated in Supplementary Table 1. Articles reporting original interventional or observational studies addressing the incidence, prevalence or risk of antipsychotic-related DIMD were included in the review. Articles unavailable in English or exclusively focusing on preclinical data were excluded. Case reports, non-systematic reviews and position/editorial papers were not cited directly unless reporting relevant data. Sample sizes and results of statistical testing demonstrating significant differences between groups in cited studies are indicated within the text when reported in original studies.

INCIDENCE OF DIMD WITH NOVEL ANTIPSYCHOTICS

A. BREXPIPRAZOLE

i. Mechanism and Unique Features

Brexipiprazole’s mechanism of action in treatment of psychosis and treatment-resistant depression (TRD) is primarily through partial agonism at D2Rs and 5-HT2A Rs with antagonism at 5-HT2C Rs [7]. Brexpiprazole has less partial agonism at the D2R relative to aripiprazole but has higher 5-HT2A antagonism which may explain its lower likelihood of inducing DIMD. Aripiprazole’s higher potency in inhibiting acetylcholinesterase activity may also play a role in its propensity to cause more DIMD relative to brexpiprazole [8]. It was approved for use in schizophrenia and as an adjunctive treatment of depression in 2015 [9].

ii. Incidence of DIMD

The most common acute DIMD associated with brexpiprazole is akathisia occurring within three weeks of drug initiation [9]. In two double-blind, randomized control trials (DBRCTs) evaluating brexpiprazole for treatment of schizophrenia (n = 636, 674), there was no significant difference in acute DIMD as measured by multiple scales, including the Simpson-Angus Scale (SAS), Barnes Akathisia Rating Scale (BARS), Abnormal Involuntary Movement Scale (AIMS), and the Drug-Induced EPS Scale (DIEPSS), at doses up to 4mg/day after 6 weeks relative to baseline or placebo [9, 10]. Other studies (n = 282, 379) have shown higher rates of akathisia relative to placebo (6.5–13.5% versus 1.0–7.1%) at doses of brexpiprazole greater than 2 mg/day [11, 12]. A six-week phase III trial (n = 459) for acute schizophrenia episodes in Japan demonstrated 9–11% of individuals in treatment groups needed medication for parkinsonism relative to 4% in the placebo group while akathisia (1.7–5.3% versus 6.9%) was less frequent in the treatment groups [13]. A 52-week open-label Japanese trial (n = 150) for maintenance treatment of schizophrenia demonstrated that the rates of akathisia and tremor to be 8.5% and 4.3%, respectively [11]. Another one-year DBRCT (n = 524) for maintenance treatment of schizophrenia demonstrated no significant increase in acute DIMD based on motor rating scales with equal rates of akathisia (1.0%) but higher rates of tremor (3.0% versus 1.0%) relative to placebo [14]. A meta-analysis of three brexpiprazole (2–4 mg/day) RCTs (n = 1444) demonstrated no significant differences in SAS, BARS, or reported DIMD compared to placebo [15].
Similarly, in two phase III DBRCTs for TRD (n = 353, 677), brexpiprazole showed minimal but significant increases in the BARS (0.18 vs 0.01 points, p = 0.0001), mixed changes in AIMS (0.08 vs 0.0 points, p = 0.014; 0.03 vs 0.04, p = 0.87) and SAS (0.18 vs −0.02, p = 0.004; 0.12 vs 0.0, p = 0.053), and no difference in DIEPSS scores relative to placebo with one individual withdrawing due to drug-induced akathisia [12, 16]. Incidence of akathisia ranged from 3–13% in these studies in a dose-dependent manner relative to placebo 1.0–2.3% [12, 16]. Other studies (n = 49–1044) have shown comparable rates of acute DIMD in both short-term and long-term studies [17, 18].

In a 52-week open-label extension of phase III trials for schizophrenia, only one of 1072 individuals developed TD, classified as moderately severe [18]. One case report demonstrated copulatory dyskinesia in association with brexpiprazole use [19]. While most data suggests that DIMD from brexpiprazole is not severe, one case report described severe parkinsonism [20]. One case report using brexpiprazole in the treatment of Parkinson’s disease psychosis demonstrated no AE [21]. Overall, akathisia is the most common DIMD with brexpiprazole but rates of DIMD in general are low.

iii. Comparison to previous SGAs
Relative to adjunctive aripiprazole use (n = 353, 677), brexpiprazole showed lower SAS and BARS scores and a lower incidence of akathisia in TRD (12, 16). In an open-label RCT for schizophrenia comparing aripiprazole (n = 33) to brexpiprazole (n = 64), DIMDs were twice as likely in the aripiprazole-treated group, most commonly akathisia (21.2% versus 9.4%), without significant differences in SAS or BARS [22]. In a small cohort of 37 Japanese individuals with schizophrenia or schizoaffective disorders, switching from other SGAs to brexpiprazole significantly improved scores by 1 point on average on the DIEPSS (p = 0.008) after eight weeks. Conversion of patients to brexpiprazole did not demonstrate any significant increase in DIMD [23]. A systematic review (n = 3925) comparing aripiprazole to brexpiprazole demonstrated no significant difference in the rates of akathisia or other DIMD [24]. In terms of akathisia, quetiapine, brexpiprazole, and ziprasidone have the highest number needed to harm (NNH) at 188, 112, and 72 in a study comparing SGAs [25]. Compared to other SGAs, brexpiprazole has a more favorable rate of DIMD in general but is prone to causing akathisia [26].

B. CARIPRAZINE
i. Mechanism and Unique Features
Relative to aripiprazole and brexpiprazole, cariprazine has a tenfold higher affinity for D3Rs with partial agonism at D2Rs and 5-HT1A Rs [27]. The D3R is preferentially expressed in brain regions, such as the ventral striatum, that regulate reward and motivation rather than motor control which could explain its proposed efficacy in the treatment of negative symptoms in schizophrenia [28]. It is FDA approved for the treatment of schizophrenia and bipolar disorder (BPD) [29].

ii. Incidence of DIMD
Safety and proof of concept trials (n = 392) with cariprazine demonstrated adverse events of total acute DIMD, tremor, and akathisia of 6–10%, 3–6%, and 9–10%, respectively [30]. A phase II DBCRT study (n = 732) in acute schizophrenia episodes for eight weeks demonstrated acute parkinsonism and akathisia in 8–10% and 11–15%, respectively, (1.5–2 times of placebo) with a 2% DIMD-related discontinuation rate [31]. Using higher doses (up to 9 mg/day) of cariprazine in a phase III trial (n = 446) demonstrated rates of acute akathisia, total DIMD, parkinsonism, use of anti-DIMD medications, and tremor of up to 17%, 10%, 17%, 24% and 5%, respectively, with no attributable discontinuation [32]. While the rates of DIMD were higher than placebo, they were only increased in the 6–9 mg/day group relative to the 3–6 mg/day group for total acute DIMD (10% versus 5%), acute parkinsonism (17% versus 9%), and use of anti-DIMD medications (24% versus 16%) [32]. Other phase III DBRCTs (n = 461) and open-label studies (n = 60) including studies up to 72 weeks have shown similar rates of acute DIMD [33, 34]. Discontinuation rates in these trials were less than 2% for acute akathisia, total DIMD, or parkinsonism [33–35, 37]. In a long-term 48-week trial (n = 679), rates of acute akathisia, parkinsonism, and total DIMD ranging from 13–23%, 11–15%, and 6.5–8.1% were observed when stratified by modal daily dose ranges [36]. Interestingly, akathisia and parkinsonism showed an inverse relationship to dose which could be due to limitations in titration based on tolerability [36]. Similarly, pooled analysis of cariprazine DBRCTs (n = 2048, 2193) for acute schizophrenia episodes using doses from 1.5–12 mg/day showed dose-dependent rates of akathisia, total DIMD, and parkinsonism of 7.8–18%, 6.0–11.3%, and 10.3–17.4%, respectively [38, 39]. Interestingly, use of cariprazine for schizophrenia in an elderly subgroup (n = 17 elderly, n = 83 total) was not associated with akathisia as an adverse event in one small study [40].

In a DBRCT (n = 819) evaluating cariprazine as an adjunctive therapy for TRD, akathisia occurred in 6.6% and 22.3% in low and high dose treatment groups compared to 2.3% in placebo [41]. Parkinsonism occurred in less than 3% of treated individuals [41]. Acute DIMD occurred in 3.0%, 6.2% and 11.0% of placebo, low and high-dose treated groups, respectively, with discontinuation in 0.4%,
0.7%, and 1.1% [41]. Use of cariprazine in trials (n = 238, 403, 233) for acute manic or mixed episodes in BPD showed similar rates of acute DIMD as studies in schizophrenia and depression [42–44]. Pooled analysis of studies using cariprazine in bipolar depression and another pooled analysis of three RCTs (n = 1065, 1407) in bipolar mania showed a dose-dependent increase in akathisia (2.2–9.6% versus 0–2.4%; NNH = 19) and other DIMD (3.0–29.2% versus 2.1–11.5%; NNH = 43) relative to placebo with discontinuation due to these AEs in 3% or less of individuals [45, 46].

In pooled analyses (n = 1407), TD was rare in the short-term, occurring in less than 0.5% of individuals over 7–9 weeks [46]. A systematic review of nine RCTs (n = 4324) in patients with BPD, schizophrenia, and MDD, cariprazine had a significantly higher risk of akathisia (relative risk [RR] = 3.92, 95% CI 2.83–5.43) and tremor (RR 2.41, 95% CI 1.53–3.79) compared to placebo [47].

iii. Comparison to previous SGAs
In a phase II trial (n = 732) for acute exacerbation of schizophrenia, cariprazine had lower acute DIMD than the risperidone (4 mg/day) group [31]. Another phase III trial (n = 461) for negative symptoms in schizophrenia and a pooled RCT analysis (n = 2193) showed rates of akathisia were slightly higher in the cariprazine group (8%) compared to the risperidone group (5%) whereas rates of total DIMD and parkinsonism were similar to slightly higher in the risperidone group (4–7% versus 6–10%) with similar rates of discontinuation (1–2%) [33, 38]. Use of 6 mg/day (n = 157) but not 3 mg/day (n = 155) cariprazine was associated with increased akathisia relative to 10 mg/day (n = 152) aripiprazole [32]. In a pooled analysis (n = 2193) of schizophrenia trials, rates of DIMD were higher for risperidone 4 mg/day (27.1%) compared to cariprazine 1.5–12mg/day (17.4%) but not aripiprazole 10 mg/day (5.9%) while akathisia rates were higher for cariprazine (11.3%) relative to risperidone (8.6%) or aripiprazole (7.2%) [38]. Rates of parkinsonism were similar in the risperidone group while half in the aripiprazole group relative to cariprazine [38]. NNH for akathisia is estimated to be ~30 for cariprazine, 34 for quetiapine, and 15 for lurasidone (n = 1407) [46]. A network meta-analysis (n = 4324) of multiple RCTs showed that there was no significant difference in overall rates of DIMD comparing aripiprazole, quetiapine, cariprazine, and lurasidone but surface under cumulative ranking curves (SUCRA) demonstrate reduced risk of DIMD for aripiprazole (overall DIMD 0.57, akathisia 0.51) and cariprazine (overall DIMD 0.45, akathisia 0.38) relative to quetiapine (overall DIMD 0.28) and lurasidone (overall DIMD 0.22, akathisia 0.13) [48]. Cariprazine may be more prone to causing akathisia, particularly early in treatment, relative to aripiprazole and risperidone while overall DIMD rates are intermediate among SGAs.

C. LUMATEPERONE
i. Mechanism and Unique Features
Lumateperone is a first-in-class medication that simultaneously acts at serotoninergic, dopaminergic, and glutamatergic receptors [49]. It was FDA approved for use in schizophrenia in 2020 but is also under investigation for mood disorders [49, 50]. At 5-HT 7 Rs, it acts as an antagonist with 60-fold higher affinity than at D2Rs, where it acts pre-synaptically as a partial agonist and postsynaptically as an antagonist, which is proposed to reduce its risk of causing DIMD [49]. By contrast, other SGAs like risperidone and olanzapine have a 12:1 affinity for binding of D2Rs and 5-HT 7 Rs. By inhibiting a serotonin transporter, it also increases phosphorylation of GluN2B receptors in mesolimbic pathways. The unique mechanism of this drug suggests that it could be useful at low doses for sleep and agitation reduction with antipsychotic and antidepressant effects at higher doses [50].

ii. Incidence of DIMD
A phase II DBRCT (n = 335) using doses of 60–120 mg/day of lumateperone in acute exacerbation of schizophrenia demonstrated AEs of akathisia in 2% of subjects, which was comparable to placebo. There were no significant change in SAS, AIMS, or BARS over 28 days [51]. Indeed, phase III DBRCTs (n = 450) and open label studies (n = 1481) with 40–60 mg/day confirm that the rates of acute DIMD are similar to placebo for as long as 12 months [52, 53]. In pooled studies (n = 381, 1073), lumateperone showed similar rates of acute DIMD and use of medications to treat DIMD compared to placebo and lower rates than risperidone (4 mg/day) [54, 55]. No clear reports of tardive syndromes associated with lumateperone have been published.

D. LURASIDONE
i. Mechanism and Unique Features
Lurasidone is approved for acute and maintenance treatment of schizophrenia and for treatment of bipolar depression [56]. It acts via potent antagonism at 5-HT 7 Rs and partial agonism at 5-HT 1 Rs mediating cognitive and antidepressant effects while having significant affinity for 5-HT 7 Rs and D2Rs [56].

ii. Incidence of DIMD
Short-term, six week, trials (n = 149, 285, 500) of lurasidone in a DBRCT for acute schizophrenia episodes showed dose-dependent total acute DIMD, parkinsonism, dystonia, and akathisia AE rates of 4–26%, 4–11%, 3–4%, and 2–24%
Changes. A meta-analysis (n = 1608) of three DBRCTs BARS, SAS, and AIMS scores did not demonstrate significant Discontinuation due to akathisia was 1% or less [66, 67].

To placebo [68].

risk ratio = 2.999; 95% CI 1.64–5.45) and for lurasidone in schizophrenia demonstrated higher rates of akathisia (8.1–14%) and parkinsonism (1.4–4.3%), dystonia (1.4–3.1%), and tremor (2.0–3.1%) [66, 67]. Discontinuation due to akathisia was 1% or less [66, 67]. BARS, SAS, and AIMS scores did not demonstrate significant changes [67]. A meta-analysis (n = 1608) of three DBRCTs for lurasidone in schizophrenia demonstrated higher rates of akathisia (risk ratio = 2.999; 95% CI 1.64–5.45) and dystonia (risk ratio = 6.013; 95% CI 1.51–23.99) compared to placebo [68].

A 24-week open-label extension trial (n = 813) for lurasidone’s adjunctive use in bipolar depression showed rates of overall DIMD and akathisia as high as 10.7% and 8.1%, respectively, with these AEs found more commonly in adjunctive therapy rather than monotherapy arms [69]. Another 28-week open label extension trial (n = 413) showed akathisia and parkinsonism in 18.2% and 7.3%, respectively [70]. Discontinuation rate of lurasidone due to akathisia was less than 3% in these studies [69, 70]. A DBRCT (n = 496) with placebo control, using a 20-week stabilization phase and subsequent 28 week treatment period in BPD showed similar rates of DIMD as other long-term studies [71]. Case reports have identified individuals with TD, drug-induced parkinsonism, rabbit syndrome, and laryngospasm with lurasidone-exposure [72–76].

iii. Comparison to previous SGAs

In a 12-month trial (n = 427) in maintenance treatment for schizophrenia, similar rates of parkinsonism (4–5%) and tremor (3%) but significantly higher rates of akathisia (14.3% versus 7.9%; NNH = 16, p < 0.05) were observed for lurasidone (40–120 mg/day) and non-significantly higher rates of dystonia for risperidone (2–6 mg/day, 3.1% versus 5.9%) [66]. Six-week trials (n = 478, 488) for acute schizophrenia episodes using extended-release quetiapine 600 mg or olanzapine 15 mg daily in a separate arm relative to lurasidone demonstrated increased rates of reported DIMD, parkinsonism, and akathisia in the lurasidone arm [77, 78]. A 12-month non-inferiority study (n = 292) comparing quetiapine XR (200–800 mg/day) to lurasidone (40–160 mg/day) for maintenance therapy in schizophrenia similarly demonstrated lower rates of DIMD (11.9–21.4% versus 3.5%) and akathisia (10.7–12.6% versus 2.4%) in the quetiapine arm [79]. A systematic review (n = 1223) of three DBRCTs comparing lurasidone, olanzapine, and quetiapine-XR in bipolar depression demonstrated significantly increased risk for quetiapine-XR causing acute DIMD (RR = 3.11, 95% CI 1.36–7.09) while both quetiapine-XR (RR = 3.71, 95% CI 1.26–10.95) and lurasidone (RR = 2.04, 95% CI 1.03–4.04) significantly increased risk for akathisia relative to placebo [80]. A short 21-day DBRCT (n = 301) comparing lurasidone (120 mg/day) to ziprasidone (160 mg/day) demonstrated comparable low rates of DIMD including akathisia [81]. In a comparative efficacy study (n = 43,049), lurasidone (OR 2.46, 95% CI 1.55–3.72) had a risk of DIMD similar to antipsychotics such as risperidone (OR = 2.09, 95% CI 1.54–2.78) and paliperidone (OR = 1.81, 95% CI 1.17–2.69) but higher than clozapine (OR = 0.3, 95% CI 0.12–0.62) relative to placebo [82]. A meta-analysis of fourteen studies (n = 6221) in bipolar depression demonstrated no difference in rates of DIMD among lurasidone, quetiapine, aripiprazole, olanzapine, and ziprasidone [83]. Lurasidone seems to have comparable risk of causing akathisia and other DIMD relative to previously marketed SGAs.

E. PIMAVANSERIN

i. Mechanism and Unique Features

Pimavanserin is a 5-HT2A-R inverse agonist and was approved for use in Parkinson’s disease (PD) psychosis (PDP) in 2016 at a dose of 34 mg daily [84]. Because of its unique selectivity
for this receptor, located on multiple types of excitatory neurons, it has benefit in psychosis without causing anti-dopaminergic side effects [85]. Decreasing signaling through the 5-HT _2A_ receptor may impact psychosis through decreasing dopamine release in the nucleus accumbens and in ventral tegmental neurons that project to frontal cortical and limbic regions [86].

**ii. Incidence of DIMD**

In placebo-controlled DBRCTs (n = 60, 199) in PDP, AEs were similar between placebo and treatment groups [86, 87]. Use of pimavanserin in conjunction with subtherapeutic doses of risperidone or haloperidol in psychosis in chronic schizophrenia did not show differences in motor AEs or as measured by SAS or BARS compared to placebo (n = 423) [84]. Very preliminary evidence (n = 403) for efficacy and similar tolerability in adjunctive treatment of negative symptoms in schizophrenia also exists without increase in DIMD while other studies are ongoing [88–91]. No studies have specifically cited or examined the risk of tardive syndromes with pimavanserin.

**iii. Comparison to previous SGAs**

No studies were identified that have evaluated the risk of DIMD in pimavanserin relative to SGAs.

**F. OLanzapine-SAMIDORPHAN (O-S)**

**i. Mechanism of action**

Olanzapine is an SGA that antagonizes 5-HT _2A/C_ Rs, 5-HT _4_ Rs, D1-4Rs, histamine-1 receptors (H1Rs) and α _1_-adrenergic receptors and has been associated with weight gain and propensity for development of metabolic syndrome [92]. Samidorphan has been shown to alter glucose homeostasis and has been used clinically to mitigate weight gain caused by olanzapine [92]. O-S was approved in 2021 for treatment of schizophrenia and BPD [92].

**ii. Incidence of DIMD**

In a four-week DBRCT (n = 401) for acute exacerbations of schizophrenia, there was no significant difference between AIMS, BARS, or SAS scores compared to olanzapine alone [93]. In the two-week follow-up, rates of parkinsonism (3.7% vs 4.5%), akathisia (6.0% versus 4.5%), and dyskinesia (1.5% vs 0.8%) were comparable between the O-S versus the olanzapine group and lower than placebo [93]. The safety profile with regard to DIMD was similar in other studies (n = 42, 234, 277 309, 561) with evaluation up to 52 weeks with no discontinuation of drug related to DIMD AEs [94–98].

**4. INCIDENCE OF DIMD WITH ALTERNATIVE FORMULATIONS OF ANTIPSYCHOTICS**

**A. LAI ANTIPSYCHOTICS AND OTHER NOVEL FORMULATIONS**

LAI antipsychotics were developed to improve poor medication adherence observed with oral formulations [99]. Studies looking at pooled data comparing oral versus LAI formulations have had mixed results with some showing no difference in DIMD whereas others have suggested higher rates with LAIs [100–102]. We review below evidence regarding incidence of acute DIMD and tardive syndromes in recently developed LAIs and other novel drug formulations.

**i. LAI Risperidone**

**1. Mechanism and Unique Features**

LAI risperidone was the first SGA available in depot form and is now available in intramuscular and subcutaneous formulations of varying dosages and half-lives [103]. LAI risperidone has lower peak levels and fluctuations in levels than oral formulations [104]. Risperidone is an antagonist at D2Rs greater than D1Rs and has inverse agonist activity at 5-HT _2A/C_ Rs [105].

**2. Incidence of DIMD**

Both short (n = 49, 264, 804) and long term (up to one year) studies (n = 215) in acute and maintenance treatment of schizophrenia and BPD have demonstrated that rates of acute DIMD, ranging from 3–23% were not significantly different between LAI risperidone and placebo and there were low not significantly different rates of discontinuation (<1%) due to DIMD [103, 106–108]. Interestingly, a 3 year study (n = 108) transitioning individuals from oral antipsychotics to LAI risperidone demonstrated reductions in TD and acute DIMD only in individuals with what was diagnosed as concurrent dopamine supersensitivity psychosis, suggesting that switching to LAIs may reduce the incidence of acute DIMD and TD in a subset of individuals with what was diagnosed as concurrent dopamine supersensitivity psychosis, suggesting that switching to LAIs may reduce the incidence of acute DIMD and TD in a subset of individuals [109]. An open-label 52-week study (n = 652) of LAI risperidone demonstrated ~1% rate of persistent TD which is comparable to oral SGAs [110]. A case of atypical TD involving the cricothyroid musculature in LAI risperidone has been reported [111]. Meta-analyses and systematic reviews comparing LAI versus oral risperidone showed no difference in DIMD [112, 113].
ii. LAI Paliperidone palmitate
1. Mechanism and Unique Features
Paliperidone, a metabolite of risperidone, has 1-, 3-, and 6-month LAI formulations [114]. Paliperidone is pharmacologically similar to risperidone with a few exceptions including less protein binding in plasma (~74%), ~60% direct renal excretion, and higher affinity for 5-HT\(_{2A/B/C/6/7}\), dopamine (D2Rs, D3Rs, and α\(_{2a/b/c}\)) receptors with dose-dependent D2R occupancy [114]. Paliperidone is pharmacologically similar to risperidone, but this effect was not significant due to high variability [130].

comparing LAI paliperidone to LAI aripiprazole showed low, comparable rates of DIMD [130].

2. Incidence of DIMD
A 12-week phase II DBRCT (n = 197) showed increased parkinsonism relative to placebo (8% versus 1%) but no other increase in DIMD [115]. A larger phase III DBRCT (n = 514) with the monthly formulation showed similar rates of DIMD, most commonly parkinsonism (5–6%), to placebo and decreased use of anti-DIMD medications from 30% before to 6% after the double-blind period but no significant change in SAS, BARS, or AIMS scales [114]. Another 12-week DBRCT (n = 305) with the 3-month formulation showed increased rates of akathisia, 4%, relative to placebo, 1%, but no difference in other rates of DIMD [116]. Other studies (n = 197–951) in schizophrenia and schizoaffective disorder with LAI paliperidone also showed low rates of DIMD [117, 118]. These results are similar to studies (n = 305, 514) with oral paliperidone and one pooled analysis (n = 3121) suggest relatively decreased DIMD rates with LAI paliperidone (anti-DIMD medication 17% versus 12%, p = 0.0035; BARS –0.09 versus −0.03, p = 0.023; SAS –0.04 versus 0.0, p < 0.0001) [114, 116, 119].

A pooled analysis (n = 3743) looking specifically at rates of TD showed low rates <0.2% regardless of administered formulation [120]. A case of unmasked parkinsonism has been reported with LAI paliperidone use persisting after drug cessation [121]. Case reports of tardive dystonia, TD, and parkinsonism, with LAI paliperidone are also found in the literature [122–124].

A 13-week open label trial (n = 452) in schizophrenia comparing LAI risperidone to LAI paliperidone showed higher rates of akathisia (19.7% versus 13%), tremor (17.9% versus 10.5%), and anti-DIMD therapy (46.2% versus 31.4%) in the risperidone group [125]. However, other RCTs and pooled analyses (n = 1214–2183) comparing LAI risperidone and LAI paliperidone showed <5% rates of DIMD without significant differences between groups [126, 127]. LAI paliperidone had comparable minimal and non-significant changes in Extrapyramidal Symptom Rating Scale (ESRS) when compared to oral SGAs in smaller RCTs (n = 84, 444) [128, 129]. One of these studies (n = 444) also suggested that LAI paliperidone had lower rates of DIMD compared to oral paliperidone [129]. Studies (n = 295) comparing LAI paliperidone to LAI aripiprazole showed low, comparable rates of DIMD [130].

2. Incidence of DIMD
Short-term studies (n = 340, 623) using LAI aripiprazole in acute schizophrenia episodes did not show significant changes in motor scales but some studies showed a higher rate of akathisia relative to placebo (11% versus 4%) [134, 135]. In a 52 week, placebo-controlled DBRCT (n = 710) of maintenance treatment of schizophrenia using LAI aripiprazole, rates of overall DIMD, akathisia, dyskinesia, dystonia, and parkinsonism were 14.9% vs 9.7%, 5.6% vs 6.0%, 0.7% vs 1.5%, 8.2% vs 3.0%, for treatment versus placebo, respectively, with no significant difference in change in AIMS, SAS, or BARS which is similar to data from other long term studies [136]. Frequency of anticholinergic therapy for DIMD in these studies was 16.7% versus 10.4% in placebo [135, 136]. A meta-analysis (n = 986) comparing tolerability of LAI versus oral aripiprazole showed no difference in DIMD [112]. A systematic review (n = 4796) comparing various LAIs demonstrated that aripiprazole had a lower odds of DIMD compared to risperidone or paliperidone, but this effect was not significant due to high variability [130]. No reports of tardive syndromes related to LAI aripiprazole were found in the literature.

iv. Transdermal asenapine
1. Mechanism and Unique Features
Asenapine is the first transdermal antipsychotic approved in 2019 by the FDA for treatment of schizophrenia. Patches are applied daily with equal hepatic and renal metabolism [137]. It antagonizes multiple receptors including 5-HT\(_{1A/1B/2B/2C/6/7}\), D1/D2/D3Rs, and α\(_{2a/b/c}\) adrenergic receptors with dose-dependent D2R occupancy [138].
III DBRCT using sublingual asenapine showed akathisia and overall DIMD rates as high as 7.7% and 10%, respectively, during a 12–16-week open label period (n = 549) and 1.6% and 2.4%, respectively, during the 26-week double-blind phase (n = 253) relative to 0.8% in the placebo group [139]. Other studies (n = 80–532) in both schizophrenia and bipolar mania or mixed episodes showed similar rates of these AEs and minimal changes in motor rating scales [138]. No reports were found of tardive syndromes are associated with transdermal asenapine.

V. Inhaled loxapine
1. Mechanism and Unique Features
Inhalation of loxapine as rescue therapy.

2. Incidence of DIMD
In clinical trials (n = 129–344) for agitation in bipolar mania and schizophrenia, inhaled loxapine was not associated with any significant increase in DIMD [141]. Benztopine was given for one individual with acute akathisia and another with acute dystonia (jaw and oculogyric) which resolved symptoms in both cases. Oral formulations of loxapine have been associated with TD, tardive dystonia, and other DIMD at rates similar to SGAs [141]. However, the risk of tardive syndrome is likely rare with intended use of inhaled loxapine as rescue therapy.

5. AGENTS WITH NOVEL MECHANISMS UNDER INVESTIGATION
Three novel classes of antipsychotics are currently under investigation whose mechanisms of action have the potential to obviate DIMD. Uloturant, a trace amine-associated receptor 1 (TAAR1) agonist, is currently in Phase 3 clinical trials for schizophrenia [142]. TAAR1s are G-protein coupled receptors expressed in multiple brain regions including the ventral tegmental area and the dorsal raphe nucleus, which allow for modulation of dopaminergic and serotonergic pathways improving positive schizophrenic symptoms and sleep [142]. While it has agonist activity at 5-HT1A Rs, it does not have antagonism at D2 or 5-HT2A Rs which is thought to improve its tolerability [142]. Early studies have not shown a risk of DIMD as expected from data in preclinical models with AEs most commonly being somnolence and GI symptoms [143].

Roluperidone is a high-affinity 5-HT1A antagonist and P2R antagonist [91]. Early trials have shown modest clinical efficacy without increased DIMD which is unsurprising given its lack of D2R antagonism [144]. Common side effects included headache, asthenia, and somnolence [91].

Finally, xanomeline is a selective M1 and M4 muscarinic receptor agonist [145]. While this drug has no significant direct binding to dopamine receptors, preclinical models demonstrate functional dopaminergic antagonism particularly in the ventral tegmental area [146]. Clinical trials have shown efficacy in schizophrenia with side effects mainly consisting of nausea, vomiting, gastrointestinal distress, and hypersalivation [145, 147]. Notably, there were no DIMD-related AEs [145, 147]. Ongoing studies will help better define the efficacy and tolerability of these medications.

6. SUMMARY
The decision to select an antipsychotic for treatment depends on an informed risk-benefit analysis of each individual patient’s condition and needs, including consideration of efficacy and a range of potential side effects. Systematic reviews and meta-analyses show that in general, FGAs and SGAs, have comparable efficacy with differences that are on a continuum rather than discrete [148, 149]. Research and development of antipsychotic drugs has largely been driven by the goal of eliminating adverse DIMD, especially irreversible tardive syndromes, while preserving antipsychotic effectiveness. The recent novel antipsychotics work through various mechanisms including greater serotonergic relative to dopaminergic antagonism, dose-dependent partial D2R agonism, and D3R antagonism [7, 27, 56]. These mechanisms allow for efficacy in psychosis while reducing incidence of DIMD (Table 2). Lumateperone and pimavanserin have a significantly lower risk of DIMD and tardive syndromes than previous antipsychotics, although relatively less data are currently available for these new agents. Relative to previous SGAs, brexipirazole has a reduced risk of overall DIMD with increased risk of akathisia. Cariprazine and lurasidone have comparable rates of DIMD relative to previous SGAs with the former having a higher relative rate of akathisia.

LAI antipsychotics have also become increasingly used in the past 10 years to improve adherence [99]. LAI paliperidone may have lower rates of DIMD whereas
| RISK OF OVERALL (ACUTE AND TARDIVE) DIMD | RISK OF TARDIVE SYNDROMES | RISK OF AKATHISIA | RISK OF PARKINSONISM | MECHANISM OF ACTION | FDA-APPROVED INDICATIONS | DRUG NAME |
|------------------------------------------|--------------------------|------------------|---------------------|---------------------|-------------------------|-----------|
| Less than previous SGAs [15, 22, 23]     | Case reported [46]       | Possibly increased relative to previous SGAs [66] | Less than previous SGAs [15, 23, 153] | Partial agonist at D2Rs and 5-HT_1A Rs, antagonist at 5-HT_2A Rs | Schizophrenia and adjunct for depression [9] | Brexpiprazole |
| Similar to previous SGAs [31]            | Less than 0.5% [31]      | Slightly increased relative to previous SGAs [33, 38, 46, 48] | Similar to previous SGAs [31] | Partial agonist at D2R, D3R, and 5-HT_1A R | Schizophrenia and BPD [29] | Cariprazine |
| No increased risk relative to placebo; lower than previous SGAs [52, 53, 55] | Not Reported | No increased risk relative to placebo; lower than previous SGAs [52, 53] | No increased risk relative to placebo; lower than previous SGAs [51-53] | Antagonist at 5-HT_1A Rs, dopamine concentration dependent antagonism at D2Rs [56] | Schizophrenia [49, 50, 56] | Lumateperone |
| Two reports (acute akathisia, acute dystonia) [141] | Not reported | Single reported individual [141] | Not reported | Postsynaptic antagonism at D2R and 5-HT_1A R [140] | Agitation in schizophrenia and mania [140] | Inhaled Loxapine |
| Similar to previous SGAs [66, 77, 78, 83] | Multiple case reports [72-76] | Similar to previous SGAs [77, 80, 82, 83] | Similar to previous SGAs [66, 77, 80, 81] | Potent antagonist at 5-HT_1A Rs, D2Rs, 5-HT_1A Rs, and partial agonist at 5-HT_2A Rs | Schizophrenia and bipolar depression [56] | Lurasidone |
| Equivalent to olanzapine [94-98]         | Equivalent to olanzapine [94-98] | Equivalent to olanzapine [94-98] | Equivalent to olanzapine [94-98] | 5-HT_1A Rs, 5-HT_2A R, D1-4R, H1R and α1-adrenergic receptor antagonist [92] | Schizophrenia and BPD [92] | Olanzapine-samidorphan |
| Equivalent to olanzapine [94-98]         | Equivalent to olanzapine [94-98] | Equivalent to olanzapine [94-98] | Equivalent to olanzapine [94-98] | 5-HT_1A Rs, 5-HT_2A R, D1-4R, H1R and α1-adrenergic receptor antagonist [92] | Schizophrenia and BPD [92] | Olanzapine-samidorphan |
| Equivalent to olanzapine [94-98]         | Equivalent to olanzapine [94-98] | Equivalent to olanzapine [94-98] | Equivalent to olanzapine [94-98] | 5-HT_1A Rs, 5-HT_2A R, D1-4R, H1R and α1-adrenergic receptor antagonist [92] | Schizophrenia and BPD [92] | Olanzapine-samidorphan |
| Not reported                              | Not reported             | Not reported     | Not reported        | 5-HT_1A R inverse agonist [86] | Parkinson’s disease psychosis [84] | Pimavanserin |
| Similar to previous SGAs and oral formulations [112, 113] | Similar to previous SGAs and oral formulations [110] | Similar to previous SGAs and oral formulations [112, 113] | Similar to previous SGAs and oral formulations [112, 113] | Antagonism at D2Rs greater than D1Rs and inverse agonist activity at 5-HT_1A Rs [105] | Schizophrenia and BPD [103] | LAI Risperidone |
| Similar to previous SGAs (possibly lower than oral formulations) [114, 116, 126] | Similar to previous SGAs [105, 120, 130] | Similar to previous SGAs [114, 116, 130] | Similar to previous SGAs [114, 116, 126, 130] | Similar to risperidone with higher affinity for 5-HT_1A R [105] | Schizophrenia and schizoaffective disorder; adjunct for BPD or major depressive disorder [114] | LAI Paliperidone |
| Similar to previous SGAs and oral formulations [112, 113, 130, 134, 135] | Similar to previous SGAs and oral formulations [112, 130, 134, 135] | Similar to previous SGAs and oral formulations [112, 130, 134, 135] | Similar to previous SGAs and oral formulations [112, 130, 134, 135] | Partial agonist at 5-HT_1A Rs and as an antagonist at 5-HT_2A Rs with a dose-dependent agonism/antagonism at D2Rs [132] | Schizophrenia, BPD, major depressive disorder [131] | LAI Aripiprazole |
| Similar to previous SGAs [138, 139]      | Similar to previous SGAs [138, 139] | Similar to previous SGAs [138, 139] | Similar to previous SGAs [138, 139] | Antagonist at multiple receptors including 5-HT_1A Rs, D1/D2/D3Rs, and α1-adrenergic receptors with dose-dependent D2R antagonism [138] | Schizophrenia [137] | Transdermal Asenapine |

Table 2 Summary of Novel Antipsychotics.
DIMD = drug-induced movement disorder, FDA = Food and Drug Administration, LAI = long-acting injectable, SGA = second-generation antipsychotic.
LAI risperidone and LAI aripiprazole have similar rates compared to oral formulations. In general, alternate formulations of antipsychotics do not seem to change risk of DIMD. In the future, current drugs in the pipeline of research and development that further spare dopamine signaling suggest antipsychotic strategies that could promise separation from the intertwined and heretofore inevitable risk of acute and tardive DIMD.

Tardive syndromes with novel antipsychotics are reported to be overall rare in the trials reviewed here. However, there are major limitations that must be considered in the interpretation of this data. Most clinical trial designs include limited follow-up, at most a year; yet TD can take years to manifest, and thus may be underestimated in typical clinical trials. Furthermore, variable prior use of antipsychotics and inconsistent washout periods of prior antipsychotics complicates

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**Supplementary Table 1** Search terms used and number of results and articles reviewed.

| SEARCH TERM                  | NUMBER OF RESULTS | EXCLUDED RESULTS | NUMBER OF ADDITIONAL ARTICLES IDENTIFIED THROUGH REFERENCE REVIEW | NUMBER OF ARTICLES SELECTED FOR REVIEW |
|------------------------------|-------------------|------------------|-----------------------------------------------------------------|----------------------------------------|
| “brexpiprazole” AND “tardive” | 7                 | 2                | 3                                                              | 10                                     |
| “brexpiprazole” AND “extrapyramidal” | 28                | 7                | 25                                                             | 46                                     |
| “cariprazine” AND “tardive”   | 5                 | 2                | 2                                                              | 5                                      |
| “cariprazine” AND “extrapyramidal” | 54                | 3                | 10                                                             | 61                                     |
| “ilumateperone” AND “tardive”  | 2                 | 0                | 5                                                              | 7                                      |
| “ilumateperone” AND “extrapyramidal” | 8                | 2                | 0                                                              | 6                                      |
| “lurasidone” AND “tardive”    | 3                 | 0                | 0                                                              | 3                                      |
| “lurasidone” AND “extrapyramidal” | 53                | 13               | 5                                                              | 45                                     |
| “pimavanserin” AND “tardive”  | 1                 | 0                | 0                                                              | 1                                      |
| “pimavanserin” AND “extrapyramidal” | 6                 | 3                | 2                                                              | 5                                      |
| “olanzapine” “samidorphan” AND “tardive” | 0                | 0                | 0                                                              | 0                                      |
| “samidorphan” AND “tardive”   | 0                 | 0                | 0                                                              | 0                                      |
| “olanzapine” “samidorphan” AND “extrapyramidal” | 0                | 0                | 0                                                              | 0                                      |
| “olanzapine” AND “extrapyramidal” | 0                 | 0                | 0                                                              | 0                                      |
| “olanzapine” “samidorphan”     | 34                | 8                | 2                                                              | 28                                     |
| “long-acting” “risperidone” AND “tardive” | 15                | 5                | 10                                                             | 20                                     |
| “long-acting” “risperidone” AND “extrapyramidal” | 106               | 31               | 0                                                              | 75                                     |
| “long-acting” “aripiprazole” AND “tardive” | 4                 | 0                | 0                                                              | 4                                      |
| “long-acting” “aripiprazole” AND “extrapyramidal” | 21                | 6                | 2                                                              | 17                                     |
| “long-acting” “paliperidone” AND “tardive” | 10                | 2                | 0                                                              | 8                                      |
| “long-acting” “paliperidone” AND “extrapyramidal” | 52                | 10               | 0                                                              | 42                                     |
| “transdermal” “asenapine” AND “tardive” | 0                 | 0                | 0                                                              | 0                                      |
| “transdermal” “asenapine” AND “extrapyramidal” | 1                 | 0                | 6                                                              | 7                                      |
| “inhaled” “loxapine” AND “extrapyramidal” | 4                 | 0                | 5                                                              | 9                                      |
| “inhaled” “loxapine” AND “tardive” | 0                 | 0                | 0                                                              | 0                                      |
determination of the prospective risk of TD and other DIMDs in both placebo and intervention arms. Variations in age and sex of participants can impact incidence of DIMD and make comparisons between studies difficult. The studies reviewed had widely variable sample sizes (from 10s to 1000s) and variable reporting of statistical comparison of adverse events. Results from larger controlled trials likely should carry greater clinical impact. The outcome scales capturing DIMDs employed vary between studies, and even the best of these, the Simpson-Angus Scale, have been criticized as insensitive compared to instruments used by movement disorders neurologists. Finally, other studies rely on the prescription of anticholinergic/antiparkinsonian medications as surrogate measures of DIMD which does not capture the underlying DIMDs.

Only further head-to-head comparison, multiyear real-world tolerability data, and assessment of DIMD with consistent neurologic examination compiled prospectively among initially drug-naïve individuals will give a more precise risk differential for these novel antipsychotics. While previous SGAs and novel antipsychotics have lower relative rates of DIMD compared to FGAs, increasing indications and widespread off-label use of SGAs and novel antipsychotics could lead to overall increased absolute prevalence of DIMD [150–152]. Understanding which of these medications have the greatest benefit-to-risk ratio is important to reduce the overall burden of DIMD.

COMPETING INTERESTS

Dr. George T. Kannarkat serves as a consultant for Guidepoint Global. Dr. Stanley Caroff served as consultant for Neurocrine Biosciences and Adamas Pharmaceuticals and received research grants from Neurocrine Biosciences and Eagle Pharmaceuticals unrelated to the current manuscript. Dr. James Morley has served as a consultant to GE Healthcare and received funding from the Department of Defense and the Department of Veteran Affairs unrelated to this review.

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