Magnetic resonance imaging of the dopamine system in schizophrenia – A scoping review

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For decades, aberrant dopamine transmission has been proposed to play a central role in schizophrenia pathophysiology. These theories are supported by human in vivo molecular imaging studies of dopamine transmission, particularly positron emission tomography. However, there are several downsides to such approaches, for example limited spatial resolution or restriction of the measurement to synaptic processes of dopaminergic neurons. To overcome these limitations and to measure complementary aspects of dopamine transmission, magnetic resonance imaging (MRI)-based approaches investigating the macrostructure, metabolism, and connectivity of dopaminergic nuclei, i.e., substantia nigra pars compacta and ventral tegmental area, can be employed. In this scoping review, we focus on four dopamine MRI methods that have been employed in patients with schizophrenia so far: neuromelanin MRI, which is thought to measure long-term dopamine function in dopaminergic nuclei; morphometric MRI, which is assumed to measure the volume of dopaminergic nuclei; diffusion MRI, which is assumed to measure fiber-based structural connectivity of dopaminergic nuclei; and resting-state blood-oxygenation-level-dependent functional MRI, which is thought to measure functional connectivity of dopaminergic nuclei based on correlated blood oxygenation fluctuations. For each method, we describe the underlying signal, outcome measures, and downsides. We present the current state of research in schizophrenia and compare it to other disorders with either similar (psychotic) symptoms, i.e., bipolar disorder and major depressive disorder, or dopaminergic abnormalities, i.e., substance use disorder and Parkinson’s disease. Finally, we discuss overarching issues and outline future research questions.

KEYWORDS
dopamine, schizophrenia, magnetic resonance imaging, substantia nigra, ventral tegmental area
Introduction

Schizophrenia is a disorder of behavior and brain that affects about 1% of the world population (1, 2). It is characterized by positive (e.g., delusions, hallucinations), negative (e.g., motivational deficits, anhedonia), and cognitive symptoms (e.g., impairments in cognitive flexibility or working memory). Schizophrenia shortens life expectancy by about 15 years and causes high socio-economic costs – for example, only around 20% of patients pursue a regular job (3–5). For decades, aberrant dopamine transmission has been proposed to play a central role in schizophrenia pathophysiology – this is called the "dopamine hypothesis of schizophrenia" (6, 7). Dopamine transmission, such as dopamine synthesis or dopamine receptor availability, is at the molecular level – part of the brain’s distributed dopamine system. The dopamine system consists of broadcasting dopaminergic neurons, their axonal projections to the rest of the brain, and modulatory synapses of their axons in target regions (Figure 1). In the human brain, the cell bodies of dopaminergic neurons are usually organized in nuclei and are distributed almost exclusively in the brainstem, particularly in the midline and rostral portion of the mesencephalon (8, 9). Substantia nigra pars compacta (SNc) and ventral tegmental area (VTA) represent the two major dopaminergic nuclei in the brain. Based on ex vivo immunohistochemical quantification in humans, around 68% of all dopaminergic neurons identified in the mesencephalon were found in the SNc and 12% in the VTA (10). The axons of dopaminergic neurons project in a topographic manner more or less to the whole forebrain, with the highest density of dopaminergic synapses being found in the striatum (11, 12). Dopaminergic synapses are mainly modulatory and are laid out like lattices in target regions (13, 14).

The earliest versions of the dopamine hypothesis of schizophrenia relied on indirect evidence: for example, the clinical efficacy of antipsychotic drugs was found to correlate with their affinity to dopamine receptors (15). More recently, direct evidence has come from human in vivo molecular imaging studies of dopamine transmission, employing particularly positron emission tomography (PET) and single-photon emission computed tomography (SPECT) (7, 12). These methods have become the gold standard for investigation of the dopamine system. They are used to study distinct aspects of transmission at dopaminergic synapses (12), for example: (i) dopamine synthesis and storage in presynaptic axon terminals, using radioactively labeled precursors of dopamine that are able to cross the blood-brain-barrier, for example 18F-DOPA; (ii) postsynaptic dopamine receptor availability, using tracers that bind to D1 or D2/3 receptors; (iii) dopamine release from the presynaptic axon terminal, by measuring the displacement of these dopamine receptor tracers after administration of amphetamine (which induces dopamine release); and (iv) availability of transporters carrying dopamine from the synaptic cleft back into the presynaptic axon terminal, using tracers that bind to dopamine transporters.

Concerning the consistency of findings using these molecular imaging methods, several meta-analyses identified increased presynaptic dopamine synthesis and storage in the dorsal striatum as the most consistent aberrance of dopamine transmission in schizophrenia (16–18). Most studies included in these meta-analyses investigated patients with significant psychotic symptoms; during psychotic remission, in contrast, striatal dopamine synthesis and storage might be decreased (19–21). For other molecular imaging measures, such as availability of dopamine receptors and transporters, no consistent alterations have been identified (16, 22, 23); also, these outcome measures are more variable in patients than in healthy subjects, hinting at possible patient subgroups (24).

While molecular imaging methods have enabled great advances in the understanding of dopaminergic pathophysiology in schizophrenia and might even predict the course of symptoms (21), there are also several downsides of these methods.

First, molecular imaging can only investigate synaptic dopaminergic transmission. Other aspects of the dopamine system, such as the structure or connectivity of dopaminergic nuclei, are not analyzed. Furthermore, since the highest density of dopaminergic synapses is located in the striatum, signal quality of molecular dopamine imaging is usually highest in the striatum, whereas signals of extrastriatal regions are often unreliable (7, 12, 25). This also applies to substantia nigra: for example, concerning substantia nigra’s presynaptic dopamine synthesis and storage in schizophrenia, results have been inconclusive so far (26–28). Second, the interpretation of molecular imaging outcome measures is still debated: for example, the outcome of 18F-DOPA PET is interpreted as presynaptic dopamine synthesis and storage. 18F-DOPA is assumed to be metabolized to 18F-dopamine by the enzyme "aromatic acid decarboxylase." However, normally, the enzyme "tyrosine hydroxylase," which acts one step before aromatic acid decarboxylase, is the rate-limiting enzyme of dopamine synthesis (29). Therefore, 18F-DOPA PET outcome measures might not adequately assess presynaptic dopamine synthesis and storage. Third, PET has limited spatial and temporal resolution. The spatial resolution, usually in the range of a few millimeters, is limited in itself by the "traveling" of positrons over several milliseconds before annihilation. The temporal resolution is...
FIGURE 1
Overview of the human dopamine system. (A) The dopamine system in humans. Schematic depiction of the human brain’s dopamine system, captured by the distribution of both dopamine cell bodies in the midbrain, i.e., in substantia nigra pars compacta (SNC) and ventral tegmental area (VTA), and dopamine D1 and D2 receptors mainly in striatum and frontal cortices (adapted with permission from 12). Note the topographic arrangement of dopamine projections, indicated by the same colors at different levels of the dopamine system. (B) Dopaminergic cells and nuclei. Top: Dopaminergic cells in the midbrain (Tyrosine hydroxylase immunostaining) [adapted with permission from (168)]. Bottom: Boundaries between the red nucleus (RN), parabrachial pigmented nucleus (PBP), substantia nigra (SNC and SNr), and subthalamic nucleus, overlaid on a T1w template of midbrain MRIs with isotropic voxel size of 700 µm [adapted from (158), open-access]. (C) Dopamine synapses and axon arborization in mice. Top left: Scheme of the modulatory nature of dopamine synapses [adapted with permission from (13)]. Bottom left: Lattice-like, volume-focused distribution of dopamine synapses in the striatum of mice [adapted with permission from (14)]. Right: Dopamine axon arborization in the striatum of mice (adapted from (169), open-access). (D) Dopamine connections of VTA. Diffusion-weighted MRI-based tractography of human VTA into the cortex (adapted with permission from (168)).

limited by the use of graphical physiological modeling, which requires steady-state conditions that usually develop only after several minutes. In 18F-DOPA PET, for example, outcome measures are currently obtained only after 70 min of scan time at minimum. Fourth, scan costs are usually rather high, both due to tracer production and long scan times. Fifth, radiation exposure impedes repeated longitudinal examinations.

For these reasons, it is necessary to complement molecular imaging with other in vivo imaging tools to map the dopamine system and its changes in schizophrenia. Magnetic resonance imaging (MRI)-based methods that investigate distinct aspects of the dopamine system are promising candidates for such tools, at least for the study of further aspects of the dopamine system beyond synaptic transmission. In the following, we perform a scoping review on MRI of the dopamine system in schizophrenia; particularly, we will focus on four MRI methods which have been applied in schizophrenia: (i) neuromelanin MRI, which is thought to measure long-term dopamine function in SNC and VTA; (ii) morphometric MRI, which is assumed to measure the volume of SNC and VTA; (iii) diffusion MRI, which is assumed to measure fiber-based structural connectivity of SNC and VTA; and (iv) blood oxygen level-dependent (BOLD) functional MRI (fMRI) at rest, which is thought to measure blood oxygenation fluctuations of SNC and VTA and their functional connectivity, i.e., their correlations with other brain regions’ ongoing blood oxygenation fluctuations.

For each of these four methods, we initially describe its underlying signal, its outcome measures, and its downsides. Then, we present the current state of research in (i) schizophrenia, (ii) other psychiatric disorders – with a focus on disorders with similar (psychotic) symptoms, e.g., bipolar disorder or major depressive disorder, or with assumed dopaminergic abnormalities, e.g., substance use disorder –, and (iii) Parkinson’s disease, which is frequently used as a model for dopamine MRI due to dopamine cell loss in this disease.

After these method-specific paragraphs, we summarize and discuss the main findings as well as methodological issues and missing points within and across modalities. Furthermore, we outline questions for future research.
Distinct magnetic resonance imaging-based approaches for studying the dopamine system in schizophrenia

Neuromelanin magnetic resonance imaging

Neuromelanin is a dark pigment produced by catecholamine metabolism. Over the life span, it gradually accumulates in the form of neuromelanin-iron complexes within organelles in the cell bodies of catecholamine-producing neurons, mostly in SNc, VTA, and locus coeruleus (Figure 2A) (30–33). It is thought to reflect a stable long-term marker of catecholamine metabolism, since it accumulates slowly and is not removed from neurons; its content in a given region only decreases after cell death, e.g., during neurodegeneration.

Since neuromelanin-iron complexes are paramagnetic (34), they shorten T1 relaxation and reduce magnetization transfer effects of nuclear magnetic resonance in tissues (Figure 2A) (35–38). These are probably the main factors contributing to neuromelanin-sensitive MRI. Neuromelanin MRI employs fast spin-echo or gradient-echo sequences, recently with direct magnetization transfer pulses, and shows neuromelanin-rich regions as hyperintense.

A typical outcome measure of neuromelanin MRI analysis is the contrast-to-noise ratio (CNR), which describes the ratio between the signal in a presumably neuromelanin-rich region, for example substantia nigra, and the signal of a reference region that presumably contains no neuromelanin, for example crus cerebri (Figure 2B) (39, 40). While neuromelanin MRI has been used to quantify dopaminergic neuron loss in neurodegenerative conditions for some time (39, 41, 42), it has recently also been validated to measure dopamine function in vivo in subjects without neurodegeneration: Cassidy et al. showed a relationship between substantia nigra CNR and both post-mortem neuromelanin concentration in the substantia nigra and amphetamine-induced striatal dopamine release measured by PET (43).

Downsides of the neuromelanin MRI signal comprise, first, the issue whether neuromelanin MRI specifically measures neuromelanin concentrations or also other substances/processes; further multimodal work combining imaging and histology is needed here. Second, although the spatial resolution of neuromelanin MRI (about 0.5 × 0.5 × 1.5 mm) is much higher than with PET, enabling the investigation of substantia nigra subregions (43), more refined investigations require improved sequences with better spatial resolution. For example, the ventral tegmental area is more difficult to image with neuromelanin MRI than the substantia nigra since it is smaller and contains less neuromelanin (43, 44). Furthermore, limited spatial resolution can lead to partial volume effects (45). Third, more work is needed to understand the temporal dynamics of neuromelanin accumulation, e.g., how fast one can expect signal changes when conditions (for example, psychotic symptoms) change. Fourth, the neuromelanin MRI signal could be confounded by many other non-structural factors, e.g., medication, about which reliable data are not yet available (40).

In patients with schizophrenia, studies have shown a reliable elevation of substantia nigra CNR (Figure 2C and Table 1), whereas for locus coeruleus CNR, no clear trend has been observed. The first study by Shibata et al. reported a significantly increased substantia nigra CNR, but no significant alteration of locus coeruleus CNR (46). A more recent study by Cassidy et al. found increased substantia nigra CNR only in patients with strong psychotic symptoms (43). The current literature is summarized by two recent meta-analyses, which showed a consistent significant increase of substantia nigra CNR, but no consistent alteration of locus coeruleus CNR (40, 47). However, medication effects could not be assessed, since too few data were available. Furthermore, there are mixed results concerning the relationship between substantia nigra CNR and psychotic symptoms: one study observed a positive correlation (43), while others found no relationship (48, 49). Finally, longitudinal data or data from specific patient groups, e.g., psychotically remitted patients, are still lacking.

Regarding other psychiatric disorders, increased substantia nigra CNR has been shown for substance use disorder (50, 51). For major depressive disorder, there are only few studies so far: one study reported reduced locus coeruleus CNR, but no change in substantia nigra (46); another study found that neuromelanin MRI can be employed to visually discriminate depressed patients from healthy subjects (48); and a recent study reported that CNR of substantia nigra and ventral tegmentum correlates with gait speed in late-life depression (52). So far, we know of no studies conducting neuromelanin MRI in bipolar disorder.

In Parkinson’s disease, neuromelanin MRI has been widely used. The first study by Sasaki et al. observed reduced CNR of both substantia nigra and locus coeruleus in patients (39). Two recent meta-analyses showed that patients with Parkinson’s disease can be diagnosed with neuromelanin MRI-based measures (e.g., intensity or volume) with a sensitivity and specificity of more than 80%, respectively (41, 42).

Magnetic resonance imaging-based morphometric analysis of dopaminergic nuclei

For the analysis of the macrostructure of dopaminergic nuclei, structural magnetic resonance imaging based on T1- or T2-weighted sequences can be used (Figure 3A). These sequences are sensitive to differences regarding longitudinal...
FIGURE 2
Neuromelanin MRI in schizophrenia. (A) Neuromelanin MRI signal. Schematic depiction of the metabolization of dopamine into neuromelanin-containing organelles and how neuromelanin influences the magnetic resonance imaging (MRI) signal. On the right, an exemplary preprocessed neuromelanin MRI image of one healthy subject is shown, with focus on the mesencephalon. (B) ROIs for neuromelanin MRI analysis. Typical regions of interest (ROIs) (43) used for contrast-to-noise-ratio analysis, which calculates the ratio between the MRI signals of substantia nigra and a reference region, e.g., crus cerebri. (C) Neuromelanin MRI changes in schizophrenia. Schematic depiction of the consistent elevation of substantia nigra contrast-to-noise-ratio in patients with schizophrenia (SCZ) compared to healthy controls (HC), based on meta-analytic evidence (40, 47).

(T1w) or transverse (T2w) relaxation time of free water magnetization across tissues (53). These signals can be used to estimate the volume of dopaminergic nuclei, which are anatomically defined using either manual delineation or automated techniques (Figure 3B). Automated techniques can be divided into normalization-based and segmentation-based approaches (54). Concerning normalization-based approaches, whole-brain voxel-based morphometry (VBM) is a common tool. VBM uses non-linear spatial normalization (based on a template image) to transform T1-weighted images into standard space (55). Subsequently, each voxel’s probability of belonging to a standard gray matter mask is computed. The outcome of VBM is interpreted as voxel-wise gray matter density or volume (depending on the exact methodology). A downside of VBM is that regions such as substantia nigra or other brainstem nuclei, which are not part of standard gray matter masks due to their low contrast in T1w MRI, cannot be reliably investigated with this method; instead, the whole brainstem is segmented as white matter (56, 57).

Concerning segmentation-based tools, FreeSurfer is a common tool. FreeSurfer defines brain regions-of-interest on the subject level based on atlases and tissue borders (58, 59). The outcome of FreeSurfer is interpreted as the volume of a region-of-interest in mm³. A downside of FreeSurfer is that regions-of-interest have to be predefined in the software framework based on an atlas. For substantia nigra, for example, no such predefined region-of-interest exists, thus it cannot be investigated.

Taken together, a reliable automated segmentation of the substantia nigra and ventral tegmental area remains challenging (60, 61). This is mainly due to the low contrast of conventional T1-weighted MRI images in the brainstem. The rather high iron content in these brainstem nuclei shortens T1 and thus leads to a brighter contrast in the images, which makes it demanding to distinguish from white matter (Figure 3C) (62). Although probabilistic anatomical atlases for the substantia nigra and ventral tegmental area exist, region of interest-based analysis cannot be performed, as midbrain and brainstem are classified as white matter structure by common morphometry toolboxes, which also makes whole-brain analysis results in these areas questionable. Furthermore, post-mortem-based definitions of the human dopaminergic VTA/SN and their delineation in standard spaces such as MNI are not available.
## TABLE 1 Summary of dopamine MRI studies in schizophrenia.

| Study | Patients/controls | Field strength | Technique | (Primary) outcome measure | Findings in patients |
|-------|------------------|----------------|-----------|--------------------------|---------------------|
| **Neuromelanin MRI** | | | | | |
| (46) | 20/34 | 3T | FSE | CNR of SNc vs. SCP decussation | Increase |
| (48) | 23/23 | 3T | FSE | CNR of SNc vs. SCP decussation | Increase |
| (173) | 52/52 | 3T | 3D-SPGR | CNR of SNc vs. midbrain tegmentum | Increase |
| (43) | 33/30 | 3T | 2D GRE-MT | CNR of SNc vs. CC | No difference (increase only in highly psychotic patients) |
| (51) | 30/8 | 3T | FSE | Contrast ratio of SN | Increase |
| **Morphometric MRI** | | | | | |
| (69) | 936/784 | 1.5T, 3T | 3D MP-RAGE, SPGR | Whole-brain SBM, VTA | Increase |
| **Diffusion MRI and structural connectivity** | | | | | |
| (93) | 24/22 | 3T | DWI; 4 b0s $b = 1,300 \, \text{s/mm}^2$, 42 directions | Probability of belonging to fiber bundle and FA for VTA to NAcc, mOFC, IOFC, DLPCF and amygdala tracts | Increased probability of belonging to fiber bundle in patients for VTA-left amygdala and VTA-left mOFC tracts No difference in FA |
| (106) | 22/23 | 3T | DWI; 8 b0s $b = 900 \, \text{s/mm}^2$, 51 directions | Tract dispersion and FA of Striato-nigro-striatal tract | Increase tract dispersion in patients. No difference in FA |
| (103) | 46/44 | 3T | DWI; 4 b0s $b = 1,300 \, \text{s/mm}^2$, 42 directions | FA of slMFB | Increase in severe psychotic affective patients compared to non-psychotic patients and controls |
| (92) | 30/24 | 3T | DWI; 1 b0 $b = 1,000 \, \text{s/mm}^2$, 12 directions | VTA/SNc Connectivity-based parcellation into limbic, prefrontal and sensorimotor parcels and evaluation to FA and streamline density index within these parcels | No difference in streamline density index nor FA in limbic, prefrontal sensorimotor parcels between groups |
| (104) | 46/35 | 3T | DWI; 1b0 and $b = 1,000 \, \text{s/mm}^2$, 32 directions | Connectivity index of VTA pathways to DLPCF, OFC, insular cortex | No difference |
| **Blood oxygenation MRI and functional connectivity** | | | | | |
| (145) | 21/21 (UM-SCZ/HC) | 3T | GRE-EPI, 3D MP-RAGE | VTA/midbrain seed-based FC | Decrease FC with precuneus, IPC, dACC, MCC, FG, IC, THAL, HPC, CAU, PU, cerebellum |
| (146) | 26/22 (M-SCZ/HC) | 3T | GRE-EPI, 3D MP-RAGE | VTA seed-based FC | Decrease FC with VLPCF, IC, LOC Increase with DLPCF |
| (147) | 71/64 (M-SCZ/UM-SCZ/no records) | 3T | GRE-EPI, 3D MP-RAGE | SN/VTA seed-based FC | Decrease with ACG, FG, CAU, OC, PAL, CAU, PU, PC/PCC, AG, TG, PG, PH, FG, OPER, CAL |

ACG, anterior cingulate gyrus; AG, angular gyrus; CAL, calcarine cortex; CAU, caudate nucleus; CC, crus cerebri; CNR, contrast-to-noise ratio; D-SCZ, deficit schizophrenia; dACC, dorsal anterior cingulate cortex; DLPCF, dorsolateral prefrontal cortex; DWI, diffusion-weighted imaging; FC, functional connectivity; IPC, inferior parietal cortex; FG, fusiform gyrus, FG, superior, middle and inferior frontal gyri; FG, superior, middle and inferior frontal gyr; GRE, gradient recalled echo; HC, healthy controls; EPI, echoplanar imaging; IC, insular cortex; LOC, lateral occipital complex; M-SCZ, medicated schizophrenia patients; MCC, middle cingulate cortex; mOFC, medial orbitofrontal cortex; IOFC, lateral orbitofrontal complex; MP-RAGE, magnetization-prepared rapid gradient-echo; MT, magnetization transfer; MTG, middle temporal gyrus; NAcc, nucleus accumbens; ND-SZ, non-deficit schizophrenia; OC, olfactory cortex; OPER, Rolandic operculum; PAL, pallidum, PC/PCC, precuneus/posterior cingulate cortex; PG, postcentral gyrus; PH, parahippocampal gyrus; PU, putamen, SC, structural connectivity; SCP, superior cerebellar peduncle; SN, substantia nigra; SNc, substantia nigra pars compacta; slMFB, supero-lateral medial forebrain bundle; TG, superior, middle and inferior temporal gyrus; THAL, thalamus; UM-SCZ, unmedicated schizophrenia patients (18 schizophrenia, 3 schizoaffective disorder); VLPCF, ventrolateral prefrontal cortex.

at the moment – in contrast to the cholinergic basal forebrain, enabling VBM-based volumetric approaches (63, 64). FSL currently released a beta version of a toolbox called MIST, which is planned to provide tools to segment brainstem nuclei based on T1-weighted images in an upcoming release (65). Another possibility to segment nuclei in the brainstem and midbrain is an online processing pipeline called voBrain, which provides automated segmentation of deep gray matter nuclei, including substantia nigra, red nucleus, and subthalamic nucleus, based on T2-weighted images (66) – however, this is performed by the authors upon request, impeding large-scale applications.

For these reasons, manual segmentation remains the gold standard to delineate dopaminergic nuclei. Downsides of this approach are: it requires high expertise, is prone to inaccuracies, and is impracticable for applications to large cohorts (60, 61).
FIGURE 3
MRI-based morphometric analysis of dopaminergic nuclei in schizophrenia. (A) T1- and T2-weighted relaxation curves. The figure shows the recovery of the longitudinal magnetization (T1 relaxation). Tissues with a short T1 relaxation time such as fat and white matter (WM) recover faster and produce a stronger signal. The T2 relaxation reflects the decay of transverse magnetization. Tissue with a short T2 relaxation decays more quickly and produces a weaker signal. (B) Anatomy of SN and VTA. Axial view of the SN (blue) and VTA (red) derived from a probabilistic atlas shown on the MNI anatomical template (158). (C) T1- and T2-weighted images showing SN and VTA. Axial T1w (left) and T2w (right) structural imaging of the brainstem. The arrows point to the area of SN and VTA. (D) Unclear morphometric changes of dopaminergic nuclei in schizophrenia. Due to the low number of studies and methodological challenges, no clear trends have emerged so far.

Further – more general – downsides of T1w-based morphometric MRI derive from its underlying signal. The MRI signal has a high sensitivity to common artifacts and non-structural factors, such as head motion, breathing effects, substance use, and body weight (67). In addition, observed morphometric changes, especially in the short term, might rely on perfusion changes instead of volume changes (68).

In schizophrenia, we are not aware of any region-of-interest-based studies that directly focused on VTA or SNC – therefore, no clear trend of alterations has emerged so far (Figure 3D and Table 1). However, a few whole-brain VBM studies reported morphometrically altered clusters that contained SNC and VTA. For example, Gupta et al. (69) investigated differences in gray matter concentration using source-based morphometry, which is a data-driven extension to VBM based on independent component analysis, and found increased gray matter density in the VTA in patients with schizophrenia. They found no association with the severity of positive or negative symptoms. Moreover, some meta-analyses of whole-brain VBM studies in schizophrenia reported altered VBM scores in brainstem clusters partly covering VTA/SNC (70, 71). For example, across 127 VBM studies including over 13,000 patients and healthy controls, we observed reduced gray matter volume in a cluster located mainly in the brainstem but partly overlapping with SNc (71). The interpretation of such findings, however, is obviously difficult since these meta-analyses were not focused on dopaminergic nuclei as regions of interest.

Concerning other psychiatric disorders, e.g., bipolar disorder, major depressive disorder, or substance use disorder, we are not aware of any direct morphometric studies of dopaminergic nuclei either. Although some voxel-based whole-brain analyses revealed a morphometric increase in substantia nigra or midbrain in bipolar and major depressive disorder, and a decrease in substance use disorder (e.g., (72–74)), coordinate-based meta-analyses did not show consistent effects in SNC or VTA across studies (75, 76).

The few studies in psychiatric diseases are in stark contrast to the research in Parkinson’s disease, where several studies about morphometric differences of the dopaminergic nuclei have been performed so far. For example, Geng et al. (77) manually segmented the substantia nigra based on proton density- and T2-weighted images and did not find any alterations in different stages of Parkinson’s disease. A meta-analysis on substantia nigra volumetry reported a significant decrease (78). However, the included studies were
heterogeneous regarding image processing and anatomical labeling of the substantia nigra.

**Diffusion magnetic resonance imaging-based analysis of structural connectivity of dopaminergic nuclei**

To date, diffusion-weighted imaging (DWI) is the only technique available to study white matter fibers and tracts in vivo in humans. DWI relies on the principle that water molecules are in constant motion. In free water (i.e., water molecules not bound to, for example, macromolecules), molecules can equally move or diffuse in all directions – this is called isotropic diffusion. In biological tissues comprising certain cell architectures, the diffusion of water molecules is hindered by cell membranes, thus resulting in anisotropic diffusion (79–81). Thus, by measuring the orientational dependence of water diffusion, it is possible to estimate the orientation of underlying membranes, for example axons, and to delineate fiber pathways (82, 83). DWI is sensitive to water diffusion by applying magnetic field gradient pulses in different directions (b-vectors; minimum 6, usually 32 to 64 in clinical settings), in order to map water diffusion not only for a spectrum of directions but also for each voxel across the whole brain (79–81). The gradients’ strength and timing is defined by the so-called b-value: the higher the b-value, the stronger the diffusion effect.

Magnitude and directionality of water diffusion can be modeled by so-called 3D diffusion tensors (DTI, diffusion tensor imaging). In mathematical terms, a voxel-wise diffusion tensor is a $3 \times 3$ matrix, which subjects to diagonalization and results in a set of three eigenvectors. Such a tensor can be visualized as an ellipsoid, for which the three eigenvectors represent the major, medium, and minor axis of the ellipsoid, respectively, and the corresponding three eigenvalues represent the apparent diffusivities along these axes (Figure 4A). From the tensor, one can estimate both the direction of maximum diffusivity and the diffusivity in any arbitrary direction (84). Several measures are derived from the diffusion tensor, such as fractional anisotropy (FA), which describes how strongly the diffusion is pronounced to one main direction (85).

While FA can inform about the microstructure of white matter, it does not provide information about pathways of white matter, for example tracts connecting two brain regions such as VTA and parts of the prefrontal cortex. However, 3D representations of fiber bundles can be obtained by identifying streamlines of adjacent primary eigenvectors of diffusion across neighboring voxels (86). This procedure is called tractography, and its output is called structural connectivity. In other words, streamlines can be defined as the adjacent sets of 3D points produced by tractography algorithms (87). There are two main types of tractography: In deterministic tractography (88), each streamline is generated following the main direction of diffusion of each voxel step by step, until a too wide curvature angle or a low signal stops the pathway. Probabilistic tractography (89), in contrast, considers the uncertainty of the main orientation of diffusion and estimates a distribution of potential orientations; next, a streamline is generated following each of these possible orientations. The higher the number of streamlines per voxel, the higher the probability for that voxel to be part of a fiber bundle (Figure 4B). Using probabilistic tractography, it is possible to parcellate seed regions according to predefined target regions, which are known to project into the seed, by the highest number of streamlines into the seed region (90, 91).

Several limitations of DWI and DTI-based tractography make the study of dopaminergic connectivity challenging. First, DWI is typically of low spatial resolution (i.e., about $2 \times 2 \times 2$ mm), which makes tractography of small regions such as VTA difficult. Therefore, some studies have opted for manually delineating these structures (92) or drawing small spheres around brain coordinates of VTA/substantia nigra with subsequent tractography from there (93, 94). Second, tractography is unspecific to the type of neuronal projections, i.e., one cannot differentiate, for example, between dopaminergic and glutamatergic connections from VTA. Third, microstructural measures of tracts such as FA are unspecific to microscopic features, reflecting a wide range of processes or structures such as myelin content, axonal packing, axonal diameter, or gliosis (95). Thus, there is a need for more specific methods, for example, myelin water imaging (96) or neurite orientation dispersion and density imaging (97). Fourth, tractography is not sensitive for the direction of tracts, efferent or afferent relative to a seed region (95, 98). Fifth, tractography is unreliable in regions where fibers have more than one main orientation, such as regions of crossing or fanning fibers (99–101). To overcome this limitation, more accurate models were developed, for example constrained spherical deconvolution (CSD), which is used to estimate the distribution of fiber orientations within each imaging voxel, regardless of the number of underlying fiber orientations (102). Finally, there is a wide range of distinct measures obtained from both DWI and DTI-based tractography, which are – confusingly – “lumped together” under the label “structural connectivity,” making comparisons across studies challenging. Examples of outcome measures comprise the number of streamlines per voxel; the percentage of streamlines starting from the seed region (for example VTA) that reach the target region (for example orbitofrontal cortex), also referred to as connectivity index; the volume of streamlines in the seed region divided by the total volume of the seed region [also called the streamline density index (92)]; or the probability for a given voxel to be part of an a priori defined tract connecting two regions (93). Changes in any of these measures for the pathway connecting VTA and orbitofrontal cortex, for example, would indicate alterations in this pathway; however, the exact underlying biological process (number of axons, degree of compaction of axons, etc.) remains
FIGURE 4
Structural connectivity of dopaminergic nuclei in schizophrenia based on diffusion-weighted MRI. (A) Principles of diffusion-weighted imaging (DWI). Water molecules are free to diffuse in all directions in free water while their diffusion is restricted in biological tissue such as axonal fiber bundles and will follow their main orientation. It can be quantified using a mathematical model called a tensor which can be represented as an ellipsoid with its main axis representing the principal direction of diffusion. In the case of free water there is no principal direction of diffusion thus the ellipsoid takes the shape of a sphere. (B) Structural connectivity outcome measures. Example of a DWI volume. All DWI volumes are used to perform tractography between two regions of interest. In deterministic tractography, each reconstructed streamline follows the main direction of diffusion per voxel or while in probabilistic tractography as represented here, a distribution of possible orientations is modeled and each streamline is reconstructed from each possible orientation. In the reconstructed path, the value in each voxel represents how many streamlines passed through it. The higher number of streamlines per voxel, the higher the probability for that voxel to be part of a fiber bundle. (C) VTA and SN structural connectivity in healthy controls (HC). Example of probabilistic tractography from VTA and SN showing their connectivity to the rest of the brain [adapted with permission from (107)]. (D) Mixed structural connectivity findings in schizophrenia. Structural connectivity can be estimated by a variety of different outcome measures, making comparisons across studies difficult; so far, no clear trends have emerged.

unclear. In addition, measures of white matter microstructure can also be referred to as structural connectivity, as is the case for FA (103, 104) or for tract dispersion index (105), an index of fiber geometry which reflects the degree to which tracts deviate from being parallel such as in fanning fibers, for example (106). As for previously described measures of structural connectivity, FA changes in the VTA-orbitofrontal cortex connection might reflect several processes such as axonal degeneration, demyelination, gliosis, or characteristics of fiber architecture such as kissing or fanning fibers as would also be indicated by the tract dispersion index.

Several studies have investigated both substantia nigra and VTA structural connectivity in healthy humans. Most studies performed either DTI-based or CSD-based probabilistic tractography with seeds in the VTA or SN, respectively (91, 107, 108). An alternative approach is, first, performing whole-brain tractography and then selecting the superolateral part of the medial forebrain bundle (MFB) which connects the VTA to both frontal lobe and forebrain (109–111) to specifically target connections between VTA and parts of the prefrontal cortex such as the orbitofrontal cortex (112). Finally, one study compared substantia nigra and VTA connectivity among each other, observing higher scores of substantia nigra connectivity for several brain regions including primary motor and somatosensory cortex, premotor cortex, and caudate nucleus (Figure 4C) (107). The most detailed study of substantia nigra connectivity to date is probably the one by Zhang and colleagues, who used probabilistic tractography and confirmed in humans a tripartite connection organization that was described before in monkeys by the use of similar techniques: the medial substantia nigra connects with limbic striatal and cortical regions, the ventral substantia nigra with associative regions of cortex and striatum, and the lateral substantia nigra with somatomotor regions of striatum and cortex (91).
Few studies have investigated VTA/substantia nigra structural connectivity in schizophrenia so far, each using a different measure of structural connectivity. Thus, findings are rather mixed, and no clear pattern has emerged so far (Figure 4D and Table 1). Bracht and colleagues (93) performed DTI-based probabilistic tractography from VTA to ventral striatum and prefrontal cortex. They found higher probability values for belonging to certain fiber bundles in schizophrenia patients compared to healthy controls in both the left VTA-to-orbitofrontal cortex and left VTA-to-left amygdala pathway. These probability values of both pathways were associated with patients’ negative symptom severity (measured by the negative sub-scale of positive and negative syndrome scale, PANSS). Basile and colleagues used CSD-based tractography to perform a connectivity-based parcelation of SN/VTA regions based on their connections to limbic, prefrontal, and sensorimotor cortices (92). Between patients and controls, they observed no difference in the parcelation into “limbic,” “prefrontal,” and “sensorimotor” territories across groups, and additionally no differences in both streamline density index and FA in these territories. Using connectivity index and FA as measures of structural connectivity between VTA and dorsolateral prefrontal cortex, orbitofrontal cortex, and insular cortices, Giordano and colleagues reported no differences between patients and controls either (104). However, two other studies, which used a whole-brain tractography approach first and then identified pathways of interests such as the striato-nigro-striatal pathway or the medial forebrain bundle including VTA/substantia nigra connections, reported mixed findings (103, 106): altered tract dispersion index was found in patients with schizophrenia (106), while the medial forebrain bundle FA was unchanged in most patient subgroups of schizophrenia spectrum except for those with severe psychotic symptoms – they had increased FA values compared to both patients with less psychotic symptoms and healthy controls (103).

In other psychiatric disorders, most studies investigated the microstructure of the medial forebrain bundle. For example, in major depressive disorder, Bracht and colleagues used a bilateral probabilistic fiber tracking approach to extract pathways between the VTA and ventral striatum and medial and lateral prefrontal cortex, respectively, and compared mean FA values (93) between patient subgroups and healthy controls. They found reduced FA only in patients during a severe depressive episode. In bipolar disorder, using a similar methodology, Denier and colleagues did not find any FA differences (113). Regarding substance use disorders, lower FA in the VTA-Nucleus accumbens tract as part of the medial forebrain bundle has been reported in individuals with stimulant use disorders (114), and lower FA in several white matter tracts has been reported in adolescents and young adults with a history of cannabis use (115) as well as in cocaine, nicotine and alcohol users (116).

In Parkinson’s disease, lower FA (117) and reduced structural connectivity concerning several measures such as streamline density were reported for nigrostriatal projections in patients with Parkinson’s disease (118, 119).

**Blood oxygenation magnetic resonance imaging of dopaminergic nuclei and their blood oxygen level-dependent resting-state functional connectivity**

Blood oxygenation reflects the amount of oxygenated hemoglobin of erythrocytes in the blood. Blood oxygenation level dependent functional magnetic resonance imaging (BOLD fMRI) is a neuroimaging technique that non-invasively depicts local changes of blood oxygenation along time (120). First, concerning physical underpinnings of the BOLD signal, deoxygenated hemoglobin has paramagnetic properties, leading to altered decay of transversal magnetization of free and hemoglobin-bound water (i.e., H + -nuclei of water molecules) under MRI conditions (T2* -weighted imaging, gradient-echo echo-planar imaging) (121, 122). Fast sampling rates enable mapping of changes in T2* relaxation of water magnetization in a certain voxel, which are then interpreted as BOLD changes along time, with temporal and spatial resolutions between 0.5–3 s and 2–3 mm, respectively (Figure 5A). Second, concerning neurophysiological and physiological underpinnings of BOLD changes, blood oxygenation is a summary measure reflecting a couple of processes, namely changes in neuronal activity and neurovascular coupling processes, which link neuronal activity with blood oxygenation. Neurovascular coupling includes hemodynamic processes (e.g., blood volume and flow changes), vascular processes (e.g., vascular reactivity based on myocyte and endothelial activity), and mediating processes (e.g., astrocytic activity) (123) (Figure 5A). In the human cortex, the BOLD signal comes mainly from perforant and branching arterioles and venules (123). In gray matter nuclei such as VTA/SNC, BOLD changes reflect changes in the neuronal and neurovascular processes mentioned above, including those directly related to dopaminergic cells and their local cellular support network. Already from this short description, one can infer that the mapping of VTA/SNC BOLD fluctuations onto dopaminergic neuronal activity is rather unspecific, weak, and limited by fMRI’s temporal and spatial resolution.

During resting states, the spontaneous BOLD fMRI signal fluctuates in an infra-slow frequency range, 0.01 to 0.15 Hz (124). The correlation between spontaneous BOLD fluctuations of different regions or voxels, e.g., between seed regions such as VTA/SNC and cortical regions of the forebrain, can be studied via pair-wise correlation coefficients (mainly normalized Pearson’s correlation coefficients) (125, 126) (Figure 5B). Correlation coefficients are interpreted as BOLD functional...
connectivity (FC) of ongoing brain activity and are typically used to investigate seed-based FC patterns, reflecting the seed's static functional connectivity architecture (124, 127) (Figures 1D, 5C). Particularly, FC patterns of VTA/SNc have been used to investigate the functional architecture of the dopaminergic system with the assumption that any BOLD fluctuation coherent (or correlated) to the VTA/SNc time course might be related to dopaminergic projections from the VTA/SNc toward, for example, the forebrain (128, 129). The resulting FC pattern in healthy participants indeed overlaps largely with patterns of tracer-based anatomical dopaminergic projections identified in primates, including projections to the ventral striatum, amygdala-hippocampal complex (130), prefrontal and cingulate cortices (130, 131), as well as thalamus (132) (Figures 1D, 5C). Furthermore, one should note that BOLD FC is indeed sensitive to external dopaminergic challenges, suggesting that changes in BOLD FC might reflect changes in dopamine transmission (133–136).

However, potential downsides of the BOLD fMRI signal, BOLD FC in general, and BOLD FC of the VTA/SNc in particular have to be mentioned. In general, BOLD fMRI is sensitive to several confounds, including, for example, scanner-related artifacts such as hardware-related instabilities and undesired variations in the magnetization field, scanning-dependent low frequency drifts, and subject-induced motion artifacts, making multi-scanner or multi-site studies challenging (137). Specifically related to VTA/SN, cardiac and respiratory-related artifacts, such as arterial pulsation, are important in the brainstem due to its proximity to large pulsing vessels (138–141). Additional challenges concern the preprocessing of images, specifically the co-registration process, where
BOLD-related T2*-weighted images with a limited spatial resolution (i.e., voxel size $\sim 2 \times 2 \times 2$ mm) are mapped to high-resolution T1-weighted structural images for anatomical reference (i.e., $\sim 1$ mm) causing possible spatial distortions, mismatched alignment, and partial volume effects in brainstem structures. Although there are several advanced scanning and preprocessing procedures accounting for these limitations (e.g., high-resolution fMRI with stronger magnetic fields such as 7T MRI, movement-induced artifact detection and application of correction algorithms) (142–144), BOLD FC patterns for VTA/SNc seeds should be evaluated carefully.

In patients with schizophrenia, studies of VTA/substantia nigra resting-state BOLD FC demonstrated an overall trend to reduced functional connectivity between VTA/substantia nigra and forebrain structures in patients (145–147). Hadley and colleagues were the first to study VTA BOLD FC in patients with schizophrenia or schizoaffective disorder. Patients were initially unmedicated and received antipsychotic treatment with risperidone during the study. They were scanned before start of the treatment, after 1 week, and after 6 weeks, while symptom severity was assessed weekly. The seed for the analysis was defined as a 3-mm spherical region of interest at Montreal Neurological Institute (MNI) coordinates $[0; 16; -7]$ representing the VTA/midbrain (148). Before the start of treatment, unmedicated patients showed reduced VTA/midbrain FC toward several cortical and subcortical regions compared to controls (145) (Figure 5D and Table 1). After 1-week treatment, connectivity toward the thalamus was restored; however, connectivity strength was not correlated to treatment response. Interestingly, pre-treatment connectivity strength toward the dorsal anterior cingulate cortex was positively correlated with a good response to treatment, whereas pre-treatment strength toward areas of the so-called “default mode network” demonstrated a negative correlation with treatment response (145). With a similar VTA definition, this pattern of widespread reduced VTA FC was largely replicated by Giordano and colleagues in medicated clinically stable patients (146), and later by Xu and colleagues using a VTA/substantia nigra seed in both medicated and unmedicated patients (147). This FC reduction was correlated with patients’ motivational deficits (146, 147). In contrast, increased resting-state BOLD FC in schizophrenia was observed only by Giordano and colleagues, namely hyperconnectivity between the VTA and the dorsolateral prefrontal cortex, interpreted as an imbalance between the so-called “salience” and “executive central” resting-state networks (146). A fourth study did not find any significant differences regarding VTA connectivity between patients and controls; around 60% of the sample were chronic patients (149).

In other psychiatric disorders, VTA/SNc resting-state BOLD FC results are mixed. For instance, regarding substance-related and addictive disorders, reduced connectivity between substantia nigra and caudate nucleus as well as between VTA and medial orbitofrontal cortex, anterior cingulate cortex, and nucleus accumbens has been observed in opioid use disorder (150), while reduced connectivity of VTA with olfactory tubercle, medial orbitofrontal cortex, and nucleus accumbens was found in patients with internet gaming disorder (42, 151). In both major depressive disorder and bipolar disorder, concerning VTA FC, studies observed reduced connectivity with frontal regions (149, 152), cerebellum and medio-dorsal thalamus (152) in patients, increased connectivity with pre- and postcentral gyrus, frontal and temporal gyrus, occipital lobe, pons (152), limbic regions (149, 152), cerebellum, and posterior cingulate cortex (149), or no changes between groups of patients and controls (153). Concerning substantia nigra FC in major depressive disorder patients, both reduced connectivity with anterior cingulate cortex and cerebellum and increased connectivity with pre- and postcentral gyrus, insula, inferior frontal gyrus, superior temporal gyrus, parahippocampus, and occipital lobe have been reported (152).

In Parkinson’s disease, the only study so far observed reduced FC of substantia nigra with thalamus, pallidum, caudate, putamen (154).

Discussion: Current research gaps and potential developments in the field

We performed a scoping review of four MRI methods imaging the dopamine system in schizophrenia, namely neuromelanin MRI, morphometric MRI, diffusion MRI, and BOLD resting-state fMRI. We set these studies in context regarding methodological underpinnings (with focus on the imaging signal and its downsides), typical results in healthy controls, and aberrances in both other psychiatric disorders frequently comprising psychotic symptoms and Parkinson’s disease, the reference disorder for MRI-based dopamine system imaging. A sufficient number of studies with homogeneous methodology (five) was identified only for neuromelanin MRI (Table 1), for which meta-analyses have already been conducted. For the other three modalities, the number of studies was limited (e.g., morphometric MRI) and/or studies were quite heterogeneous in terms of outcome measures or patient groups (e.g., diffusion MRI) (Table 1). In the following, we first give an overview of the included methods and what aspects of the dopamine system they measure; second, we discuss findings of dopamine MRI in schizophrenia for each imaging modality regarding homogeneity, signal downsides, findings in other disorders, and relation to symptoms; third, we outline missing points and limitations across modalities; finally, we outline some questions for future research.
Summary of modalities and their coverage of the dopamine system

While molecular imaging, e.g., PET, of the dopamine system in schizophrenia focuses on aberrant dopamine transmission at the synaptic level, MRI-based imaging methods complementarily focus on other parts of the dopamine system (Figure 1): neuromelanin MRI focuses on cumulative catecholamine metabolism in dopaminergic nuclei, morphometric MRI on the structure of dopaminergic nuclei, diffusion MRI on structural tract-based connections of dopaminergic nuclei, and BOLD resting-state fMRI on functional blood oxygennation-based connectivity of dopaminergic nuclei. Thus, the four methods investigate different but complementary aspects of the dopaminergic system. Neuromelanin and morphometric MRI measure localized processes within dopaminergic nuclei, while BOLD fMRI measures signal coherence between dopaminergic nuclei and other brain regions. Diffusion MRI, on the other hand, can measure microstructure within dopaminergic nuclei, but also the macro- and microscopic architecture of fibers/tracts in regions rather distant from dopaminergic nuclei, which presumably contain fibers from/to dopaminergic nuclei.

Findings in schizophrenia for each modality, comparison to other disorders, and problems

Neuromelanin MRI studies showed a consistent increase of contrast-to-noise-ratio in the substantia nigra in patients with schizophrenia (40, 47) (Figure 2). This is interpreted as increased accumulation of neuromelanin in the substantia nigra, which is an indicator of increased long-term catecholamine metabolism. The methodology across studies was rather homogeneous – only the reference regions for contrast-to-noise ratios varied. Downsides of the signal comprise limited knowledge about the specificity of the signal for neuromelanin, influences from factors such as medication, limited spatial resolution, and the very low temporal resolution (presumably in the range of years). Concerning substantia nigra contrast-to-noise ratio in other disorders, there is a similar trend for increase in substance use disorder, no clear trend in major depressive disorder or bipolar disorder, and a clear trend for reduction in Parkinson's disease. There is no clear trend for a link with psychotic symptoms; longitudinal studies are lacking.

The few morphometric MRI studies of dopaminergic nuclei showed no clear trend for morphometric alterations in SNc and VTA in patients with schizophrenia (Figure 3): while the study of Gupta and colleagues (69) demonstrated increased volume in the brainstem overlapping with SNc, using whole-brain VBM in a large patient sample (＞900), the meta-analysis of whole-brain VBM studies in schizophrenia by Brandl et al. (71) found reduced VBM scores in a similar brainstem cluster. As a major limitation of these approaches, it has to be noted that we identified no study directly focusing on dopaminergic nuclei (e.g., with a ROI-based approach); instead, we only identified whole-brain studies/meta-analyses of T1-based morphometry that observed changes overlapping with dopaminergic nuclei. Downsides of the signal include the unclear biological underpinnings (particularly regarding the temporal dynamics of the signal), no specificity for dopaminergic neurons, low contrast in dopaminergic nuclei, which impedes automatic delineation and necessitates manual segmentation with ensuing problems, as well as sensitivity to artifacts. Concerning morphometric MRI measures of dopaminergic nuclei in other disorders, there is a trend for increase in major depressive disorder and bipolar disorder, a trend for decrease in substance use disorder, and consistent decrease in Parkinson's disease. In schizophrenia, the relationship with symptoms is unclear. We know of no longitudinal studies in schizophrenia.

Diffusion MRI studies in schizophrenia showed alterations in several structural connectivity measures, hinting at an aberrant macroscopic fiber architecture and fiber microstructure. However, due to the considerable heterogeneity of methods and measures, no clear pattern has emerged so far (92, 93, 103, 104, 106) (Figure 4). Further downsides of the signal include low spatial resolution, no specificity regarding dopaminergic neurons, unclear underlying microscopic processes, inability to determine the directionality of fiber tracts, and limited reliability in areas with diffuse fiber orientations. Concerning structural connectivity of dopaminergic nuclei in other disorders, there is a trend for reduction in major depressive disorder and substance use disorder, no clear trend in bipolar disorder, and a trend for reduction in Parkinson's disease. In schizophrenia, despite inconsistent findings, at least one study suggested stronger aberrances in patients with more psychotic symptoms. We are not aware of any longitudinal studies.

Resting-state BOLD fMRI studies in schizophrenia showed a trend for reduced resting-state functional connectivity of dopaminergic nuclei with widespread forebrain areas (145–147) (Figure 5). However, results were heterogeneous – some studies reported no group differences or even increased functional connectivity --, making conclusions difficult. Downsides of the signal include the lack of specificity for dopaminergic neurons, influences from non-neuronal sources, limited spatial resolution, sensitivity to artifacts (e.g., motion), and varying processing methods. In schizophrenia, the link to symptoms or longitudinal changes have not been investigated so far, to our knowledge. Concerning functional connectivity of dopaminergic nuclei with various subcortical and cortical regions in other disorders, both decreases and increases were observed in major depressive disorder and bipolar disorder, and a trend for reduction in substance use disorder and
Parkinson’s disease. In schizophrenia, there are hints at links between dopaminergic nuclei functional connectivity at rest and both positive and negative symptoms of schizophrenia. One longitudinal study showed a correlation between resting-state functional connectivity alterations and treatment response.

**Overarching problems across modalities**

First, neither morphometric MRI nor diffusion MRI nor BOLD fMRI are specific for dopaminergic neurons. In contrast, neuromelanin MRI is assumed to be rather specific for catecholaminergic neurons, and with sufficiently accurate delineation of dopaminergic nuclei, it can be assumed to have good specificity for dopaminergic neurons. To test and potentially increase the specificity of dopamine MRI modalities, multimodal studies are necessary, combining dopamine MRI modalities with either dopamine PET, the gold standard for *in vivo* dopamine imaging, – for example in the form of simultaneous PET/MRI – or neuromelanin MRI, as the most specific dopamine MRI approach (Figure 6). To our knowledge, only one multimodal study combined neuromelanin MRI with dopamine release PET (and additionally with arterial spin labeling-based perfusion MRI) in a combined sample of healthy subjects and patients with schizophrenia (43). They reported a link between neuromelanin contrast-to-noise-ratio and dopamine release in the associative striatum; however, no effect of schizophrenia on this link was found. In Parkinson’s disease, on the other hand, a recent study reported a patient-control difference concerning the link between neuromelanin MRI and dopamine synthesis and storage, measured by 18F-DOPA PET (155). Further systematic multimodal studies in patients with schizophrenia are needed, also encompassing other modalities. Naturally, such approaches are only correlative, i.e., they cannot determine whether observed correlations are direct or mediated via further processes. If no correlations were to be observed, on the other hand, this would not preclude links with other aspects of the dopaminergic system currently not accessible to dopamine MRI.

Second, the number of studies in general and the level of systematic investigations in particular was low. For example, we identified only one study of morphometric MRI in dopaminergic nuclei, whereas there is a large number of studies in Parkinson’s disease, which shows the general feasibility of this approach. Regarding diffusion MRI, on the other hand, there is a plethora of different outcome measures, which makes comparisons very difficult. Furthermore, more studies with a precise clinical characterization of patient samples, e.g., regarding symptom status or medication, are needed. Most studies included in this review reported only limited patient characteristics. For example, the current disease state is usually not sufficiently reported – e.g., are patients currently in a psychotic episode or in remission of psychotic symptoms? Future studies need to characterize investigated patients more thoroughly. Longitudinal approaches could clarify how measures change over time and with changing symptom status, as has been done for other MRI measures [e.g., for thalamocortical structural connectivity see (156)]. For neuromelanin MRI, for example, Cassidy et al. found no difference in substantia nigra contrast-to-noise-ratio between patients at high risk for psychosis and patients with diagnosed schizophrenia (43). Such approaches should be extended to longitudinal measurements within the same patients with schizophrenia, e.g., from psychotic phases to psychotic remission. Moreover, it is incompletely understood how dopamine MRI measures are influenced by antipsychotic drugs, whose clinical effects greatly rely on their blockade of dopamine receptors. More studies comparing medicated and unmedicated patients are necessary. Finally, also studies contrasting treatment-responsive with treatment-resistant patients should be conducted, to further elucidate mechanisms of resistance to antipsychotic drugs. This has already been done in multimodal MRI-PET studies, for example combining 18F-DOPA PET-based striatal dopamine synthesis capacity with prefrontal gray matter volume (157) or frontalstriatal functional connectivity (21). Future studies should also apply these approaches to MRI methods focusing on dopaminergic midbrain nuclei.

Third, it remains challenging to accurately delineate/segment the dopaminergic nuclei, which is necessary for all presented modalities, particularly for those without any specificity for dopaminergic neurons (i.e., morphometric, diffusion, and BOLD MRI). As already described further above, automated segmentation of SNc/VTA is not reliably feasible so far due to insufficient contrast properties of standard MRI techniques. Thus, typical publicly available masks are based on manual delineation of structures in “standard samples” (43, 158, 159). No MRI atlas of dopaminergic sub-nuclei (e.g., substantia nigra subfields) exists so far, as opposed to the cholinergic system (63, 160), for example, where a post-mortem based probabilistic cholinergic basal forebrain delineation in standard brain space is available. Furthermore, accurate delineation requires sufficient spatial resolution. The spatial resolution of the presented MRI techniques is usually not lower than 0.5–1 mm. Given the size of dopaminergic nuclei [VTA ∼60 mm³ (161); SN ∼350 mm³ (162)], this resolution appears sufficient for a satisfactory delineation of nuclei – still keeping in mind partial volume effects –, but a reliable investigation of sub-nuclei is not possible so far. New techniques like 7T MRI might alleviate these problems in the future, as suggested by promising recent attempts – however, so far, they also rely on manual delineation (163). Furthermore, the multimodal combination with neuromelanin MRI with its good contrast might improve delineation (Figure 6). Finally, instead of using neuromelanin MRI, T1w, or T2w
images for delineation, quantitative susceptibility mapping (QSM) could be used for delineation and also morphometric analysis. This novel technique provides enhanced contrast for substantia nigra and VTA by quantifying iron content based on reconstructing magnetic susceptibility sources (164). A recent study manually segmented the substantia nigra based on quantitative susceptibility maps and reported a volume increase in first-episode schizophrenia patients (150).

Future research questions

First, as already outlined above, integrative multimodal approaches are needed, combining different dopamine MRI methods to investigate complementary aspects of the dopamine systems. This could also improve the delineation of dopaminergic nuclei. Moreover, comparing dopamine MRI methods to dopamine PET methods would help to validate the specificity and reliability of outcomes (Figure 6).

Second, multimodal approaches could be extended to translational approaches, e.g., in rodents, and usage of human post-mortem data. Such studies would strengthen our understanding of the link between dopamine MRI and dopaminergic cellular processes. One example is the study by Cassidy et al., who validated neuromelanin MRI as a proxy measure of dopamine function by combining neuromelanin MRI with neurochemical measurements of neuromelanin concentration in post-mortem tissue of healthy subjects (43). Similar approaches could be conceived for other modalities.

Third, as outlined above, longitudinal approaches in patients with schizophrenia should test whether and how dopamine MRI measures change over time and with changing symptom status, e.g., from psychotic phases to psychotic remission. This would further enhance our understanding of dopaminergic pathology in different phases of schizophrenia and could contribute to the refinement of antidopaminergic medication strategies.

Fourth, schizophrenia is characterized by remarkable heterogeneity, not only at the symptom level, but also on genetic and pathophysiological levels (165, 166). Multimodal approaches on the dopamine system in schizophrenia might help to address the question of heterogenous changes in schizophrenia for the dopamine system, i.e., whether there are subgroups that are particularly affected in the dopamine system or in certain parts of it.
Fifth, there are cases of patients with schizophrenia in whom failures of the dopamine system do not appear to dominate (e.g., patients resistant to antidopaminergic medication). As alternative underpinning pathophysiology, other disturbed transmitter systems might be relevant, e.g., glutamate or acetylcholine (7). Furthermore, there might be additional pathophysiology not dominated by transmitter system disturbances, e.g., aberrant neurovascular units (167). Therefore, future studies and reviews should also deal with these questions.

Conclusion

Although the number of studies is still low, neuromelanin MRI, morphometric MRI, diffusion MRI, and BOLD resting-state fMRI have proven to be viable and complementary methods to investigate the human dopamine system in vivo in schizophrenia. We observed a clear trend of results only in neuromelanin MRI – due to a sufficient number of studies and homogenous outcome measures –, suggesting increased neuromelanin accumulation in the substantia nigra in patients with schizophrenia. A clear trend for gray matter volume changes in VTA/SNC morphometry is not observable so far due to the low number of studies and methodological challenges. In diffusion MRI, there was no clear trend either, largely due to the heterogeneity of outcome measures used across studies. In resting-state BOLD fMRI, there was a weak trend for reduced resting-state functional connectivity of dopaminergic nuclei with forebrain regions, whose interpretation is difficult since the direction of BOLD changes cannot easily be interpreted. As major overarching problems across modalities, we identified most modalities’ lack of specificity for dopaminergic neurons, the low number of studies – particularly systematic investigations of specific patient groups and longitudinal studies are lacking –, and the accurate segmentation of dopaminergic nuclei. Advanced novel MRI techniques and multimodal studies, particularly linking dopamine PET and MRI, might alleviate these problems in the future.

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All authors contributed to conceiving, researching, and writing of this manuscript.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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