INTRODUCTION

Poststroke aphasia is a common language disorder that occurs after cortical damage and white matter (WM) disconnections in dorsal (sound-to-articulation) and ventral (sound-to-meaning) streams in the left hemisphere [1]. Communication hinges, however, on more than words. It requires the ability to extract meaning from prosody, that is, the melody and rhythm in speech. Prosody lends structure...
to the speech stream by marking boundaries, signaling emphasis, or determining lexical meanings (e.g., the difference between “greenhouse” and “green house”). Moreover, prosody conveys emotional states of the speaker. These functions, respectively, are classically subsumed under the terms linguistic and affective prosody [2]. Perturbations of prosody perception (aprosodia) [3,4] affect approximately 30% of right hemisphere stroke patients [5] and have a strong negative impact on patients’ social relationships [6] and well-being [7].

Linguistic and affective prosody perception has been proposed to rely on dorsal and ventral streams in the right hemisphere [8,9], that is, frontotemporal regions interconnected dorsally via the arcuate fasciculus (AF) and ventrally via the inferior fronto-occipital fasciculus (IFOF). These streams stand in dynamic exchange with the left hemisphere language networks via the corpus callosum (CC) [10–12]. Although prosody perception deficits have been associated with right-hemispheric damage [4,13,14], especially with frontotemporal cortical structures belonging to the ventral stream [15,16], only little is known about the role of WM damage in aprosodia [17]. Modern lesion studies endorsing a hodological approach to prosodic deficits are lacking [18].

Defective processing of rhythmic–melodic acoustic patterns also occurs in the musical domain—in poststroke amusia, a common deficit affecting up to 60% of acute stroke patients [19,20]. Like aprosodia, amusia has been associated with right ventral stream damage [21–23], including perturbations of the IFOF. This suggests coincident impairments in prosody and music perception. However, comparative studies evaluating a potentially shared neuroanatomical basis of aprosodia and amusia are lacking.

Here, we evaluate and compare damaged neural structures underlying deficits in prosody and music perception using lesion–symptom mapping [24] and diffusion tensor imaging (DTI), at subacute and 3-month stages in a sample of 39 stroke patients. Based on small-scale lesion studies on prosodic deficits [4,13–15,25], lateralization of affective speech processing in the healthy brain [26,27] and previous evidence on amusia [21–23], we hypothesize that both disabilities arise from overlapping lesions in the right hemisphere, particularly from damage and disconnection of the right ventral stream.

**MATERIALS AND METHODS**

**Subjects and study design**

We present data of 39 patients (17 female and 22 male, mean age = 56.5 years, SD = 14.7) hospitalized between 2013 and 2015 at the Neurocenter, Turku University Hospital for an acute ischemic stroke (n = 28) or intracerebral hemorrhage (n = 11) in the left (n = 21) or right hemisphere (n = 18) with subsequent cognitive and motor deficits. Inclusion criteria were acute unilateral stroke, right-handedness, <80 years of age, capability to communicate in Finnish, residence in southwest Finland, ability to cooperate, and normal hearing.

Patients with prior neurological or psychiatric disease or substance abuse were not included. Most patients (82%) had a stroke within the middle cerebral artery territory, and 18% had a stroke within the posterior cerebral artery territory, with a mean lesion size of 50.1 ml (SD = 50.3). Written informed consent in accordance with the Declaration of Helsinki was acquired from all patients, and the protocol was approved by the Ethics Committee of the Hospital District of Southwest Finland. No ethnic data were collected. All patients underwent magnetic resonance imaging (MRI) and neuropsychological assessments within 3 weeks after stroke onset (mean = 12.1 days, SD = 5.5) and 3 months poststroke (mean = 100 days, SD = 8.8). Two time points were used to ascertain the stability of the findings. All patients received standard care and rehabilitation for stroke. There were no missing data.

**Assessment of prosody and music perception**

Prosody perception was evaluated with two well-validated tasks. In the linguistic prosody task [28], 30 utterances with different prosodic word stress patterns were played to the patients via headphones. Word stress denoted either a compound word or a phrase composed of two words, for example, “KISsankello” and “KISsan KELLo” (comparable to English “BLUEbell” referring to the name of a flower and “BLUE BELL” referring to a colored ringing object). After each utterance, patients were asked to select via button press one of two pictures on a screen matching what they had heard.

In the affective prosody task [29], patients were presented with 96 one-word utterances (“Saara”, a female first name) spoken with happy, sad, angry, afraid, surprised, or neutral prosody. Patients were asked to select which of the six emotions displayed on screen was expressed by pressing one of six buttons.

Music perception was assessed with a shortened version [30] of the Montreal Battery of Evaluation of Amusia (MBEA) [31]. Both subtests comprise 14 pairs of short piano melodies, half of which are identical and half of which contain a musically altered tone in the second melody. Patients were asked to judge on each trial whether the two melodies sounded identical. In the Scale subtest, the altered tones have an out-of-scale pitch change. In the Rhythm subtest, the alteration is a change in the duration values of two adjacent tones in the melody. MBEA Scale and Rhythm subtests were used separately as indices of musical pitch and rhythm perception, respectively.

Scores on both prosody and music tests at both time points were converted to percentage-correct scores and used in the analyses.

According to the established cutoff values of the MBEA, 21 patients were amusical at the subacute stage, and 16 at the 3-month stage. For the two tests used to assess prosody perception, no clear cutoff values have been established. However, comparable neurologically healthy listeners have a mean score of 83%, with SD = 9% [28]. Following the cutoff values for MBEA, patients scoring 2 SD below the healthy listeners were classified as aprosodic (i.e., <65%; mean score of the two prosody tests). At the subacute and 3-month stages, 24 and 16 patients were classified as aprosodic, respectively.
MRI data acquisition and preprocessing

Patients were scanned on a 3-T Siemens Magnetom Verio scanner at the Department of Radiology of Turku University Hospital. T1-weighted anatomical scans (flip angle = 9°, repetition time [TR] = 2300 ms, echo time [TE] = 2.98 ms, voxel size = 1.0 x 1.0 x 1.0 mm³) were acquired as well as diffusion MRI scans (TR = 11,700 ms, TE = 88 ms, acquisition matrix = 112 x 112, 66 axial slices, voxel size = 2.0 x 2.0 x 2.0 mm³) with one non-diffusion-weighted volume and 64 diffusion-weighted volumes (b-value = 1000 s/mm²).

T1 images were preprocessed using a previously reported pipeline [21,23] using Statistical Parametric Mapping (SPM8) under MATLAB v8.4.0. Unified Segmentation with medium regularization and cost function masking was applied to achieve accurate segmentation and optimal normalization in stroke patients with lesioned brain tissue. Lesion tracing was performed manually and separately for each time point by the first author (A.J.S.) using MRICron (https://www.nitrc.org/projects/mricron). The segmented T1 images were modulated, normalized to Montreal Neurological Institute (MNI) space. Lastly, the binary lesion masks were registered to MNI space.

Diffusion MRI data were processed using the FMRIB Software Library (FSL v5.0.8, www.fmrib.ox.ac.uk/fsl). First, eddy current distortions and head motions were corrected followed by gradient matrix rotation using FSL's fdt rotate bvecs. Then, brain extraction was performed using the Brain Extraction Tool. The diffusion tensors were reconstructed using the linear least-squares algorithm included in Diffusion Toolkit v0.6.2.2 (www.trackvis.org/dtk).

Voxel-based lesion–symptom mapping

Lesion–symptom relations were evaluated with multivariate lesion–symptom mapping using support vector regression (SVR-LSM) with SVR-LSM GUI [24,32]. Eight separate SVR-LSM analyses were carried out using linguistic prosody, affective prosody, MBEA Scale, and MBEA Rhythm scores at subacute and 3-month stages. All voxels damaged in at least 10% of the patients were included in the statistical analysis. SVR-β-value maps were generated using 1000 permutations, catalogued on a voxelwise basis, and thresholded at p < 0.005. Multiple comparisons were accounted for at a familywise error rate of p < 0.00625 at the cluster level (Bonferroni-corrected). Lesion volume was controlled regressing it from both lesion and behavioral data [24,32]. Brain areas were labeled based on the Automated Anatomical Labelling Atlas (http://www.alivelearn.net/xjview).

Diffusion MRI: Deterministic tractography

The DTI analyses focused on the AF, IFOF, CC, and tapetum, as these tracts have been implicated in both prosody perception and amusia [8,10,11,15,19,23]. Furthermore, based on our study hypotheses and the expected right-lateralization of results, right inferior longitudinal fasciculus and uncinate fasciculus were also dissected. All tracts were dissected manually using deterministic tractography in TrackVis (v0.6.0.1). AF and IFOF were dissected in both hemispheres. AF was dissected in its three segments: the long segment connecting frontal and temporal lobe, the anterior segment connecting frontal and parietal lobe, and the posterior segment connecting temporal and parietal lobe. For further details, please see Sihvonen et al. [23].

Volume and mean fractional anisotropy (FA) of each dissected tract were extracted using a MATLAB toolbox [33] and imported into IBM SPSS Statistics 24. We tested which parameters of which tracts explained performance in the prosody and music tasks using eight stepwise regression analyses, one for each of the four behavioral scores at the subacute and 3-month stages as dependent variable. The two WM tract parameters of each of the 12 dissected tracts served as 24 independent variables in all models. Alpha level on each model was set to 0.00625 (Bonferroni-corrected).

RESULTS

Behavioral deficits in prosody and music perception

First, the behavioral relationship of prosody and music perception was evaluated using two-tailed Pearson correlations (Bonferroni-corrected). These showed significant positive correlations between (i) linguistic prosody and affective prosody (subacute: r = 0.71, p < 0.001; 3-month: r = 0.53, p < 0.001), (ii) linguistic prosody and music perception (MBEA Scale subacute: r = 0.62, p < 0.001; 3-month: r = 0.54, p < 0.001; MBEA Rhythm subacute: r = 0.63, p < 0.001, 3-month: r = 0.58, p < 0.001), and (iii) affective prosody and music perception (MBEA Scale subacute: r = 0.66, p < 0.001; 3-month: r = 0.62, p < 0.001; MBEA Rhythm subacute: r = 0.59, p < 0.001; 3-month: r = 0.59, p < 0.001).

Lesion patterns associated with poor prosody and music perception

Lesion–symptom mapping revealed exclusively right-hemispheric lesion patterns including frontoinsular and striatal areas for both prosodic deficits and amusia. First, lesion patterns comprising right insula or basal ganglia were associated with poor linguistic (subacute and 3-month stages) and affective prosody perception (3-month stage; Table 1, Figure 1a,b). At the subacute stage, the lesion pattern associated with linguistic prosodic deficit was centered on frontal WM and extended further into right Rolandic operculum and limbic structures (Table 1, Figure 1a).

Second, pitch and rhythm amusia were also associated with lesion patterns comprising right frontal, insular, and basal ganglia areas (Table 1, Figure 1c,d), closely resembling the regions associated with prosodic deficits. Beyond the lesion pattern shared by pitch and rhythm amusia, subacute stage pitch amusia spread more ventrally, reaching the temporal lobe and limbic regions (Table 1, Figure 1c).
The right-lateralization of poor prosody and music perception was also mirrored in the direct comparison of performance after left- and right-hemisphere stroke. At both stages, right-hemispheric patients had significantly lower linguistic prosody (subacute: $t[37] = 2.579, p = 0.014$; 3-month: $t[37] = 2.344, p = 0.025$), MBEA Scale (subacute: $t[37] = 3.971, p < 0.001$; 3-month: $t[37] = 3.840, p < 0.001$), and MBEA Rhythm scores (subacute: $t[37] = 3.101, p = 0.004$; 3-month: $t[37] = 2.482, p = 0.018$) than left-hemispheric patients. Affective prosody only reached significance 3 months poststroke (subacute: $p = 0.072$; 3-month: $t[37] = 2.540, p = 0.015$).

**WM pathways associated with poor prosody and music perception**

Following hodological views of brain function [18], we then examined parameters of relevant frontotemporal WM tracts as predictors of patients’ performance in prosody and music tasks. In all but one task (subacute rhythm perception) and consistently across both stages, weaker performance was explained by damage of the right IFOF, denoted by smaller volume or lower FA (mean $R^2$ change = 27% across seven models). This consistent involvement of the right IFOF further supports the crucial necessity of intact right ventral connectivity for normal prosody and music perception (Table 2, Figure 2). In addition to the right IFOF, FA of the CC and volume of the right uncinate fasciculus explained variance in affective prosody perception at the subacute stage and 3-month stage, respectively. Furthermore, tract parameters of the right AF were related to pitch perception at both stages. Volume of the right AF (long segment) was the main predictor of rhythm perception at the subacute stage, whereas disconnection of right IFOF and left AF (and preserved connectivity of left IFOF) explained rhythm perception at the 3-month stage.

**DISCUSSION**

The present multimodal neuroimaging study identified and compared lesion patterns and WM disconnections underlying deficits in prosody and music perception in a sample of 39 subacute stroke patients followed up at 3 months poststroke. Our main findings were (i) that both linguistic and affective prosodic deficits are explained by disconnections of the right IFOF together with lesions along the right ventral stream, and (ii) that similar lesion configurations also give rise to both pitch and rhythm amusia. Our findings argue for a frequent behavioral and anatomical coupling of poststroke aprosodia and amusia. This comorbidity has important implications for patients’ well-being and rehabilitation success.

**Right ventral stream damage underlies prosodic deficits**

Poor linguistic and affective prosody perception was associated with right frontoinsular and basal ganglia lesions. Moreover, damage to the right IFOF was the strongest predictor of both prosodic deficits at both time points studied. Poor affective prosodic perception at the 3-month stage was additionally associated with damage to the right uncinate fasciculus. No consistent involvement of left hemisphere structures or of the right AF was found. These combined data highlight the damage and disconnection of right IFOF, and associated areas, as the most likely causes of poststroke deficits in prosody perception. The right ventral stream has been suggested to play a critical role in prosody perception [8,15,16], but larger DTI studies evaluating the necessity of WM tracts have been lacking [17]. The right IFOF interconnects right frontal, temporal, and inferior parietal/occipital areas and is a major anatomical ventral stream pathway [34,35]. Its cortical termination points are known to be active during both linguistic and affective prosody perception [36]. Affective

<table>
<thead>
<tr>
<th>Condition</th>
<th>Hemisphere &amp; region</th>
<th>Subacute stage</th>
<th>3-month stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linguistic aprosodia</td>
<td>R frontal</td>
<td>IFOF, ROP, insula</td>
<td>Insula</td>
</tr>
<tr>
<td></td>
<td>R basal ganglia</td>
<td>Put, Pall, Caud</td>
<td>Put, Pall</td>
</tr>
<tr>
<td></td>
<td>R limbic</td>
<td>Amy, Thal</td>
<td>—</td>
</tr>
<tr>
<td>Affective aprosodia</td>
<td>R basal ganglia</td>
<td>—</td>
<td>Put, Pall</td>
</tr>
<tr>
<td>Pitch amusia</td>
<td>R frontal</td>
<td>IFOF, ROP, insula</td>
<td>IFG, PreCG, insula</td>
</tr>
<tr>
<td></td>
<td>R temporal</td>
<td>STG, TTG</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>R basal ganglia</td>
<td>Put, Caud, Pall</td>
<td>Put, Pall, Caud</td>
</tr>
<tr>
<td></td>
<td>R limbic</td>
<td>Amy, Thal</td>
<td>—</td>
</tr>
<tr>
<td>Rhythm amusia</td>
<td>R frontal</td>
<td>ROP, insula</td>
<td>Insula</td>
</tr>
<tr>
<td></td>
<td>R basal ganglia</td>
<td>Put</td>
<td>Put, Pall</td>
</tr>
</tbody>
</table>

Note: All results are thresholded at voxelwise $p < 0.005$ and clusterwise family wise error rate $p < 0.00625$. Abbreviations: Amy, amygdala; Caud, caudate; IFG, inferior frontal gyrus; Pall, globus pallidum; PreCG, precentral gyrus; Put, putamen; R, right; ROP, Rolandic operculum; STG, superior temporal gyrus; Thal, thalamus; TTG, transverse temporal gyrus.

**TABLE 1** Lesion–symptom mapping results
prosody perception additionally engages the right temporal pole, at which the right uncinate terminates.

The lesion patterns reported here map onto stream models of prosody perception [8,16] according to which the brain encodes prosodic information in (right) superior temporal regions [15] and integrates this information over time along the posterior-to-anterior axis of the temporal lobe [37], before cognitively evaluating its emotional or linguistic significance in inferior frontal regions [9,16]. Our data suggest that the disconnection of frontotemporal regions due to right IFOF lesions (and to a lesser degree right uncinate fasciculus lesions) hinders this prosodic information flow and, hence, gives rise to prosodic deficits. Moreover, severe enough damage to either of the frontotemporal termination territories of the right IFOF can also disrupt this processing chain, resulting in comparable deficits [34].

A recent study on affective aprosodia with a focus on right ventral stream regions of interest [15] reported right posterior superior temporal and amygdala lesions, but not frontal damage, as the best predictors of affective aprosodia, in line with earlier small-scale lesion studies [3]. Notably, these findings and our results are not mutually exclusive under the present hodological view; lesions in temporal termination regions of IFOF [35] can lead to similar disconnections and deficits as observed in the present study. Likewise, it cannot be excluded that these previous results emerge from damage to the right IFOF. Moreover, the lack of
<table>
<thead>
<tr>
<th>Model</th>
<th>Variable</th>
<th>Beta</th>
<th>T</th>
<th>p</th>
<th>F(df)</th>
<th>R²</th>
<th>R² change</th>
<th>Model</th>
<th>Variable</th>
<th>Beta</th>
<th>T</th>
<th>p</th>
<th>F(df)</th>
<th>R²</th>
<th>R² change</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R IFOF vol.</td>
<td>0.471</td>
<td>3.245</td>
<td>0.002</td>
<td>F(1, 37) = 10.528</td>
<td>0.222</td>
<td></td>
<td>1</td>
<td>R IFOF FA</td>
<td>0.537</td>
<td>3.875</td>
<td>&lt;0.001</td>
<td>F(1, 37) = 15.019</td>
<td>0.289</td>
<td>0.289</td>
</tr>
<tr>
<td></td>
<td>CC FA</td>
<td>0.462</td>
<td>3.167</td>
<td>0.003</td>
<td>F(1, 37) = 10.031</td>
<td>0.213</td>
<td></td>
<td>1</td>
<td>R IFOF vol.</td>
<td>0.617</td>
<td>4.769</td>
<td>&lt;0.001</td>
<td>F(1, 37) = 22.746</td>
<td>0.381</td>
<td>0.381</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2</td>
<td>CC FA</td>
<td>0.418</td>
<td>3.050</td>
<td>&lt;0.001</td>
<td>F(2, 36) = 9.043</td>
<td>0.334</td>
<td>0.121</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>R IFOF vol.</td>
<td>0.351</td>
<td>2.559</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>R Uncinate vol.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Note:** All models are significant at $p < 0.00625$.

**Abbreviations:** AF, arcuate fasciculus; Ant, anterior; Beta, standardized regression coefficient; CC, corpus callosum; F(df), F-value (degrees of freedom); FA, fractional anisotropy; IFOF, inferior frontooccipital fasciculus; L, left; R, right; $R^2$, $R$-squared (unadjusted); T, t-value; vol., volume.
frontal involvement in the previous study [15] may be explained by the limited number of patients with frontal lesions, and the lack of posterior temporal or amygdala involvement in the present study could be partly due to our strict statistical thresholding to control for multiple comparisons, highlighting only the most robustly involved areas.

Additionally, right basal ganglia were associated with both types of prosodic deficits, highlighting the role of subcortical brain regions in prosody perception, in line with previous studies arguing for a role of the basal ganglia in sequencing and sensory–cognitive integration of auditory prosodic information [38,39]. Studying patients with lesions restricted to the basal ganglia, yet sparing the IFOF, could shed further light on the specific roles of these regions in prosody perception.

There has been a longstanding debate on the lateralization of linguistic and/or affective prosody perception, depending on different (linguistic/affective) function or shared acoustic cues [13,14,36]. Notably, in the present study, all lesions related to poor prosody (and music) perception were right-lateralized. Overall, the present data support the proposal that fundamental acoustic dimensions of prosody (and music) as well as affective information are processed in the right hemisphere [13,14]. However, the right hemisphere stands in dynamic exchange with left-lateralized language networks via the CC [10–12], complementing and

FIGURE 2 White matter tracts associated with prosody and music perception. White matter tracts whose parameters significantly predicted prosody (linguistic, affective) or music perception (pitch, rhythm) at the subacute and 3-month stages are displayed. Boxes specify the results for tasks and stages (Table 2). seg., segment
refining prosody perception by higher level linguistic processes [40]. Accordingly, the linguistic complexity of stimuli as well as the type of task [41] could influence the lateralization. Recent evidence suggests that prosodic information processed initially in the right hemisphere is fed to domain-general emotion processing areas and integrated with semantic information, resulting in bilateral engagement [42], and that the level of left-hemispheric activations increases when the verbal complexity increases [43]. The present data indicate the relevance of the CC in affective prosody perception at the subacute stage. Future large-scale studies with other materials and more complex tasks are needed to further explore the lateralization of prosodic deficits.

**Neuroanatomical overlap of music and prosodic deficits**

The present lesion profiles of amusia are in line with those observed in previous studies [21,22]. Poststroke amusia was associated with right frontotriastal lesion patterns and damage of the right IFOF, similar to prosodic deficits. Importantly, this anatomical overlap was accompanied by closely related behavioral deficits in both prosody and music perception, indicating a high degree of shared neural networks underlying the two disabilities.

Both prosody and music perception involve encoding, integration, and evaluation of pitch sequences and temporal patterns in the fleeting acoustic signal. Accordingly, prosody and music may both trigger partly similar processing mechanisms along the ventral auditory stream [44], so that disconnection of the right IFOF and/or damage of its frontal termination territories would lead to comorbid prosodic and amusical impairments, as observed in the present study. Notably, a similar reasoning also applies to the basal ganglia that play a key role in rhythm perception [45] and auditory sequence processing [38,46] hence constituting another region, the damage of which can entail comorbid deficits in prosody and music perception. The close neuroanatomical and behavioral association of prosodic and musical impairment is in line with previous single case lesion studies [47], and also with prosody–music interactions in healthy adults [28].

Musical deficits showed one notable distinction from prosodic deficits. Lesion in the right AF predicted performance in both music tasks but was unrelated to prosody perception. Moreover, lesions in the left AF predicted deficits in rhythm perception. The relevance of AF for music is in line with previous findings implicating the dorsal stream in music perception [37]. Moreover, preserved right and increased left frontoparietal functional connectivity has been associated with amusia recovery [48]. Moreover, previous studies have implicated the dorsal stream in rhythm perception in music [49] but also in rhythm benefits for language [50]. The present data suggest no significant role for AF in perception of prosody, at least not at the word level tested here. Its relevance for more complex sentence-level prosody [11] remains a topic for future research.

**Clinical considerations**

Both aprosodia and amusia are rarely diagnosed in standard care [5] although they are relatively common poststroke deficits [5,19]. Both disorders obviously affect successful poststroke rehabilitation by hampering communication and limiting the implementation of music-based interventions and are apt to reduce patients’ quality of life by affecting social interaction and psychological well-being [7].

Accurate communication between the patient and health care personnel is crucial to ensure fluent care and rehabilitation in the stroke unit. This is particularly true in severe stroke, where early and intensive inpatient rehabilitation is recommended to achieve optimal functional gains [51]. Communication may then fail to meet patients’ needs of emotional support, especially at the acute stage, when patients are commonly depressed or confused. Aprosodia may, hence, sustain the patient’s fright and anxiety, and impede early rehabilitation. In practice, emergency doctors and nursing staff should be made aware of the potential presence of aprosodia after right hemisphere damage.

Both aprosodia and amusia are likely to limit the positive effects of music-based rehabilitation strategies, which have recently emerged as promising and inexpensive stroke rehabilitation tools [52] included in the current American Heart Association stroke rehabilitation guideline [51]. Musical interventions, for example, those included in aphasia therapy, might require a personalized format for patients with aprosodia or amusia. Given that the processing of vocal music is relatively intact in amusia [48], singing-based rehabilitation methods seem promising in amusic [19] and aprosodic [53] patients.

When proceeding toward long-term, outpatient rehabilitation, both disorders may still significantly dilute obtaining of rehabilitation goals and reduce quality of life. Especially the loss of affective communication has been associated with reduced marital satisfaction [6] and enhanced caregiver burden [15], because the patient seemingly neglects the spouse's emotions. Due to concomitant anosognosia, common in right-hemispheric lesions, the patient him/her-self may not be aware of any difficulties in emotional communication.

Amusia is likely to disrupt musical leisure activities known to mitigate depression and enhance well-being during stressful periods of life [54]. Musical activities in groups, particularly choir singing, are potential means of social integration for recovering stroke patients. Although detailed studies on the psychological effects of amusia are still pending, both aprosodia and amusia are likely to increase the risk of social isolation, reduce quality of life, and thereby subject the patient to relapsing or worsening poststroke depression, which is well known to adversely affect long-term outcome [51].

In conclusion, the present results elucidate the shared neural bases of aprosodia and amusia. In clinical practice, stroke patients with right ventral stream damage should be assessed for prosodic and musical perception deficits, preferably by a speech–language pathologist and a music therapist, due to imminent profound effects on communication and patient well-being.
ACKNOWLEDGMENTS
We express our gratitude to the staff of the Tyks Neurocenter, and the patients and their families for their participation. We thank Professors Mari Tervaniemi and Riitta Parkkola, Jani Saunavaara, and radiographers Ulla Anttalainen, Riku Luoto, and Tuija Vahtera. This study was supported by the Academy of Finland (grants 257077, 277693, 299044), Finnish Cultural Foundation (grant 191230), Orion Research Foundation, and Signe and Ane Gyllenberg Foundation.

CONFLICT OF INTEREST
The authors have no conflicts of interest to declare.

REFERENCES


ORCID
Aleksi J. Sihvonen https://orcid.org/0000-0002-4501-7338

How to cite this article: Sihvonen AJ, Sammler D, Ripollés P, et al. Right ventral stream damage underlies both poststroke aprosodia and amusia. *Eur J Neurol.* 2021;00:1-10. [https://doi.org/10.1111/ene.15148]