How to demix Alzheimer-type and PSP-type tau lesions out of their mixture -hybrid approach to dissect comorbidity-

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Abstract

Neurofibrillary tangles (NFTs), are shared between progressive supranuclear palsy (PSP) and Alzheimer disease (AD). Histological distinction of PSP and AD is possible based on the distribution of NFTs. However, neuropathologists may encounter diagnostic difficulty with comorbidity of PSP and AD. In this study, we tried to circumvent this difficulty by analyzing five autopsied brains harboring both PSP and AD pathology. Tau-positive lesions were sorted based on their cell type (neuron versus glia), and tau isoforms: three-repeat (3R) versus four-repeat (4R) tau. 16 regions were selected to map these lesions throughout the brain. 4R-tau lesions were present in all areas examined. Among them, 3R-tau lesions were absent in some areas. These 4R-selective (4R+/3R-) areas dictate prototypic distribution of PSP, not usually found in AD, such as pontine nucleus, red nucleus, inferior olivary nucleus, dentate nucleus, globus pallidus and putamen, each contained both glial and neuronal lesions. In contrast, additional 3R-tau lesions were found in hippocampal formation to neocortex, where 3R immunoreactivity (IR) was predominant over the 4R counterpart mainly in neurons as found in AD but not in PSP. Although tau lesions in central grey matter, substantia nigra and locus coeruleus are found in both AD and PSP, 4R-selectivity with glial component suggests PSP origin. Even if the presence of 3 R IR in these areas suggests AD pathology, it does not exclude the involvement of PSP-type lesion because distinction of 4R IR into PSP or AD is not yet possible. Further demixing may be possible if biochemical difference of 4R tau between PSP and AD is identified.

Keywords: Tau cytopathology, AD, PSP, Comorbidity, Isoform, Glia

Introduction

Alzheimer disease (AD) and progressive supranuclear palsy (PSP) are characterized by deposition of tau in the brain. Initially, histological definitions were based on the disease-specific distribution of argyrophilic neurofibrillary tangles (NFTs), which are quite distinct between AD [1] and PSP [31]. Another histological hallmark of PSP is tuft-shaped astrocytes (TAs) [15], which are essentially not found in pure AD. Because clinicopathological spectra of PSP and of AD are still expanding [10, 30], accurate clinical diagnosis is more and more complex and difficult. This diagnostic difficulty is much enhanced in aged population, where comorbid pathologies, such as AD or Lewy pathology, may be encountered. [10, 30]. Such comorbidity is challenging to neuropathologists; how to discriminate different types of pathology in the same brain. This is particularly problematic when a brain harbors AD-type pathology and PSP-type pathology because both are characterized by tau deposits [3, 11]. Furthermore, it is not yet known whether these two types of pathology are independent or mutually related. In this study, we selected autopsy samples carrying histological diagnoses of both AD and PSP. We tried to extract disease-specific features from these brains with AD and PSP for possible discrimination, based on the distribution of TAs and that of NFTs and immunohistochemistry for phosphorylated tau (AT8) [24], three-repeat (3R) and four-repeat (4R) tau [9]. This hybrid approach was quite successful in discriminating most, but
not all, of tau-positive lesions, suggesting that AD-type pathology and PSP-type pathology are independent without significant interaction even when coexistent in the same brain.

Patients and methods
Among one hundred and eighty autopsy cases archived at Laboratory of Structural Neuropathology, Tokyo Metropolitan Institute of Medical Science from 1999 to 2013, we picked up 5 cases (3 cases from Nitobe Memorial Nakano General Hospital and 2 cases from Yokufukai hospital) with comorbid pathologies with PSP [15] and AD [5]. Written consent from the patient's family was obtained at autopsy and this study was approved at the ethics review committee of the Tokyo Metropolitan Institute of Medical Science (authorization number 16–25).

Demographic data are summarized in Table 1. Median age was 85 years (range 80–94 years) and male female ratio was 4:1. Brain weight was 1260 g on average (range 1205–1395 g), Braak NFT stage [5] had median V (range II–VI), and Braak Amyloid stage [5] had median B (range 0–C).

Sixteen regions, selected to map PSP-specific or AD-specific lesions [2, 4–6, 14, 18, 22, 34, 40] throughout the brain include primary motor cortex (PC), putamen (PU), external segment of globus pallidus (GPE), internal segment of globus pallidus (GPI), subthalamic nucleus (STN), hippocampal formation (HF), substantia nigra

### Table 1: Demographic data on 5 patients with neuropathological diagnosis of AD and PSP.

| Case | Age at death | Sex | Clinical diagnosis | Dementia | Parkinsonism | Duration of illness (year) | Neuropathological findings | Brain weight (g) | Braak NFT stage | Braak Amyloid stage |
|------|--------------|-----|--------------------|----------|--------------|---------------------------|-----------------------------|------------------|----------------|------------------|
| 1    | 86           | F   | CHF                | –        | –            | N. A                      | PSP, AD                     | 1205             | III             | 0                |
| 2    | 94           | M   | possible DLB       | +        | +            | 3                         | LB pathology, PSP, AD, AGD  | 1260             | III             | A                |
| 3    | 81           | M   | AD                 | +        | –            | 9                         | PSP, AD                     | 1230             | V               | C                |
| 4    | 85           | M   | pneumonia          | N. A     | N. A         | N. A                      | PSP, AD, CAA                | 1395             | V               | C                |
| 5    | 80           | M   | possible PSP       | +        | +            | 8                         | PSP, AD, AGD                | 1210             | VI              | C                |

AD: Alzheimer disease, AG: argyrophilic grain disease, CAA: cerebral amyloid angiopathy, CHF: congestive heart failure, DLB: dementia with Lewy body, N. A: not available, NFT: neurofibrillary tangle, PSP: progressive supranuclear palsy

### Table 2: Regional distribution of AT8 positive lesions.

| Case | AT8 | NFT |
|------|-----|-----|
| 1    |     |     |
| 2    |     |     |
| 3    |     |     |
| 4    |     |     |
| 5    |     |     |

See Abbreviations after the text
(SN), red nucleus (RN), central grey matter (CGM), tegmentum of midbrain (mainly superior colliculi, excluding CGM; M-TEG), locus coerulescens (LC), pontine nucleus (PN) and tegmentum (P-TEG), inferior olivary nucleus (ION) and dentate nucleus (DN). Six micron-thick sections were obtained from the formalin-fixed, paraffin-embedded blocks from these 16 regions. Deparaffinized sections were subjected to Hematoxylin-Eosin (HE) stain, Klüver-Barrera (KB) stain, Gallyas silver impregnation and Campbell silver impregnation [36]. Isoform-specific antibodies directed against 3R or 4R tau [9] were used [37, 41]. Briefly, deparaffinized sections were treated for 15 min with 0.25% potassium permanganate (KMnO₄), for 3 min with 2% oxalic acid (OA), for 30 min with > 99% formic acid (FA) and for 20 min autoclaved at 121 °C in 0.05 M citrate buffer [16, 38]. After intrinsic peroxidases were inactivated by 1% hydrogen peroxide (H₂O₂) for 15

Fig. 1 Separation of tau-positive lesions into progressive supranuclear palsy (PSP) and Alzheimer disease (AD) based on their distribution and cytopathology. Tau-positive lesions of PSP shown in green (left column, a-e), typically include primary motor cortex (PC), putamen (PU), globus pallidus (GP), subthalamic nucleus (STN), central grey matter (CGM), substantia nigra (SN), red nucleus (RN), tegmentum (M-TEG), locus coerulescens (LC), pontine nucleus (PN) and tegmentum (P-TEG), inferior olivary nucleus (ION) and dentate nucleus (DN) in the cerebellum. Tau-positive lesions of Alzheimer disease (AD), shown in grey (right column A-C), are more restricted to CGM, SN and LC in the brainstem while more extended in the hippocampal formation (HF) and cerebral cortex (CC). In the five comorbid cases with PSP and AD (PSP + AD in the mid column), these tau-positive lesions are partly overlapping. Regions with neurofibrillary tangles (NFTs) with tuft-shaped astrocytes (TAS) are labeled in italics, which replicate of PSP-type distribution (left column). Those with NFTs without TAS are labeled in Bold face (HF, LC and PN), which replicate AD-type distribution. Cerebral left hemisphere (coronal), b: Midbrain (axial), c: Pons (axial), d: Medulla oblongata (axial), e: Cerebellum.
sections were incubated first with 5% horse serum in 0.01 M phosphate-buffered saline containing 0.03% polyoxyethylene (10) octylephenyl ether (Triton X-100, Wako, Tokyo, Japan; PBST). They were then incubated with either 3R tau-specific antibody (RD3 1:3000, Merck Millipore, Germany) or 4R tau-specific antibody (RD4 1:1000, Merck Millipore, Germany) diluted in the same buffer for 2 days at 4 °C [9]. They were then incubated with biotinylated secondary antibody against mouse IgG (1:1000, ABC Elite, Vector, Burlingame, CA) diluted in the same buffer for 2 h at room temperature. They were then incubated with avidin-biotin-peroxidase complex (1:1000, ABC Elite, Vector) for 1 h and visualized with diaminobenzidine and nickel ammonium chloride [38]. 4R or 3R immunoreactivity (IR) in each of 16 regions was separately evaluated semiquantitatively as none: 0, mild (1 to 5): +; moderate (6 to 10): ++; severe (>10): ++++, (lesions/visual field with ×20 objective).

In some areas, where both 3R and 4R tau immunoreactivities (IR) were coexistent, double immunofluorolabeling was performed as described previously. The 6-μm thick sections were deparaffinized for double immunofluorolabeling with antibodies against isoform-specific anti-4R tau antibody (rabbit polyclonal, Cosmo Bio Co, Tokyo, Japan), raised against amino acids 275–291 of human 4R tau, which is deaminated at N279 [8], and the anti-3R tau antibody (RD3) [9]. Sections were washed with PBST, blocked for 30 min in 5% normal goat serum/0.05% sodium azide/PBS and incubated with the polyclonal anti-4R-tau antibody (1:3000) and RD3 (1:300), diluted in the blocking buffer at 4 °C for 4 days. To reduce autofluorescence of lipofuscin, sections were treated with Sudan Black B [28]. These primary antibodies were labeled with Alexa 488 conjugated with anti-rabbit IgG (Molecular Probes, Oregon, USA, 1:200) and Alexa 568 conjugated with anti-mouse IgG (Molecular Probes, Oregon, USA, 1:200), respectively, diluted in PBS with 0.03% Triton X-100 overnight in the dark. Sections were mounted with buffered glycerol containing 0.1% p-phenylenediamine. Fluorescent signals were separately captured on a confocal system (Leica SP8; Leica Microsystems GmbH, Heidelberg, Germany) through a 63 x objective (NA 1.45).

Results

Tau-positive lesions detected by AT8 were sorted into TA and NFT and semiquantitatively mapped in 16 regions as shown in Table 2. The distribution of tau-positive lesions in these 5 comorbid cases was schematized in Fig. 1 (center column). Comparison with that of PSP (Fig. 1, left column) and that of AD (Fig. 1, right column) demonstrated that the overall tau distribution in these five comorbid cases (Fig. 1, center column) was compatible with the summation of PSP (Fig. 1, left column) and AD (Fig. 1, right column). However, distribution of TA and that of NFT were not

| TA | 4R | 3R |
|----|----|----|
| Case1 | 2 | 3 | 4 | 5 | Case1 | 2 | 3 | 4 | 5 |
| 4R | + | ++ | ++ | +++ | +++ | +++ | +++ | +++ | +++ | +++ | +++ |
| 3R | - | - | - | - | - | - | - | - | - | - | - |

Table 3 Regional distribution of 3R tau lesions and 4R tau lesions.

| 3R | 4R |
|----|----|
| Case1 | 2 | 3 | 4 | 5 |
| 4R | + | ++ | ++ | +++ | +++ | +++ | +++ | +++ | +++ | +++ | +++ |
| 3R | - | - | - | - | - | - | - | - | - | - | - |

green: 4R tau region; yellow: 3+4R tau region.
similar. Both TA and NFT were detected in RN, ION, DN, GPI, GPE and PU (Tab. 2, italics), where tau-positive lesions are rare in pure AD. These regions containing both TA and NFT replicated the distribution of pure PSP (Fig. 1, left column), suggesting that these PSP-like tau lesions are not influenced by the copresence of AD pathology (Fig. 1, right column). However, both TA and NFT were found in PC, M-TEG, CGM, P-TEG and SN, where tau-positive lesions were found in pure AD as well (Tab. 2, Fig. 1).

To distinguish PSP-type and AD-type tau pathologies in these comorbid cases, tau-positive lesions, already sorted into NFT/TA (Tab. 2, Fig. 1), were further distinguished by 3R and 4R IR as shown with their relative amount in Table 3. Their immunohistochemical profiles are displayed in Fig. 2, where 4R-selective regions (Fig. 2, a-l, left: containing 4R+/3R- tau lesions) are contrasted with 4R+3R regions (Fig. 2, m-x right: containing both 3R and 4R-positive lesions). TAs were positive only for 4R tau and found in every area examined except for LC, HF, PN. 3R tau-positive TAs were absent even when 3R-positive NFT pathology was prominent as in LC (Fig. 2w).

4R-selective regions were GPI, RN, PN, ION, DN (Fig. 2 a-l, Table 3, regions in green box), where NFTs were positive only for 4R (Fig. 2 a-l, left), while STN...
contained a few NFTs positive for 3R. In contrast, NFTs positive for 4R and 3R tau were found in 4R + 3R regions, including PC (Case 5 only), HF, CGM, SN, LC, RPN, M-TEG and P-TEG. (Fig. 2 m-x, Tab. 3, regions in yellow box). Among these 4R + 3R regions, HF and LC were characterized by dominant 3R IR in NFTs over 4R IR and lack of 4R-positive glia (Tab. 3). NFTs in PC of case 5 (Braak NFT stage VI) exhibited dominant 3R IR over 4R IR. In other 4R + 3R regions (CGM, RPN, SN, STN, M-TEG, P-TEG), 3R IR and 4R IR were comparable and tau-positive astrocytes were present in variable number. Double immunofluorolabeling (Fig. 3) for 4R tau (green) and 3R tau (red) demonstrated expected dominance of 4R tau (green) in 4R-selective regions as SN (Fig. 3a). In contrast, the proportion of 3R tau (red) and 4R tau (green) was variable from a neuron to another (Fig. 3b) in 4R + 3R regions as in CGM.

Discussion

Tau pathology of PSP-type and that of AD-type have been described separately [2, 4–6, 14, 22, 40], each representing separate entities. In this study with 5 cases harboring both PSP-type and AD-type tau pathology, we tried to discriminate PSP-type and AD-type pathologies in each brain along different criteria (distribution, participation of glial changes and tau isoforms). As initial description of PSP was based only on NFTs [31], comparison of their distribution provided a solid framework to distinguish pure PSP from pure AD (Fig. 1, Tab. 2). 4R-selective regions, characterized by the copresence of TA and NFT, both devoid of 3R tau IR (lower half of Tab. 3, in green box), include typical distribution of pure PSP (PN, RN, ION, DN, GPI, GPE and PU) [15], which are not severely affected in pure AD. In contrast, involvement of 3R tau is seen in HF and LC in this series of cases with both disorders, which strongly suggests AD-type pathology [17, 21, 33, 37] rather than PSP pathology. This assumption is reinforced by the paucity of glial involvement in these regions as in pure AD cases [22] and by the predominance of 3R tau over 4R tau (Tab. 3) [12]. Therefore, neuron-selective involvement with preferential 3R tau over 4R tau may represent AD-type pathology [3, 23] even in this comorbid series, which replicated AD-type distribution of NFT (HF, IC, LC, Fig. 4, right upper rectangle labeled AD). This is in contrast with PSP-type pathology with 4R-selective tau in both TA and NFTs (Fig. 4, left lower area in green) [11, 15], which replicated the PSP-type distribution. This operational sorting through our hybrid approach was powerful enough to distinguish origin of most of tau lesions into either AD-type or PSP-type (Fig. 4), suggesting that AD-type pathology and PSP-type pathology are independent even when these two processes are occurring in the same brain [7, 18, 19, 25, 27].

This sharp distinction of AD-type pathology and PSP-type pathology in the same brain suggests that extension of AD-type pathology and PSP-type pathology are independently guided by distinct cytopathological mechanism along disease-specific patterns without crossover even if 4R tau is found in AD and PSP. If 4R tau lesions extended transsynaptically (in a prion-like manner? [26]), AD-type and PSP-type lesions that are found together in some regions should also be found associated in their areas of projection. Because coexistent PSP-type pathology and AD-type pathology were distinct in the human brain, it is hard to explain how transsynaptic extension of 4R tau, for example if any, exhibit different types of tau pathology in the same brain. Still, there remain some ambiguities in some regions such as STN, SN, M-TEG and P-TEG (area in broken line, Fig. 4), where very small amount of 3R-positive NFTs are sometimes present. In these regions, 3R-positive NFTs (arrows in Fig. 3b, red) may be of AD-type even they are positive also for 4R tau (empty arrowhead in Fig. 3b, empty arrowhead in Fig. 3b).
However, it is not yet clear whether 4R-positive neurons without 3R tau in these areas are of AD-type or PSP-type (Fig. 3b, arrowhead, green) [17, 37], because it is not possible to distinguish 4R tau of AD-type or of PSP-type. Indeed, 4R-tau lesions of AD and those of PSP similarly exhibit argyrophilia with Gallyas silver impregnation [36] and 4R tau IR with 4R-specific antibodies such as monoclonal antibody RD4 [9] or polyclonal antibodies to 4R tau [8]. It has been reported that pretangle neurons are positive for 4R tau but not for 3R tau in both AD and PSP [17, 37]. However, this selective 4R tau IR at early phase of tau deposition is gradually replaced with 3R tau IR during evolution NFTs in AD brain [12, 39]. This is in sharp contrast with PSP or CBD brain, where 4R tau IR remains persistent without involvement of 3R even after tau-positive fibrils are dense enough to form aggregated inclusions [20, 32]. Because currently available antibodies against 4R tau immunolabel tau deposits both in AD and in PSP/CBD brains [8], it remains to be clarified how 4R tau in AD brain and that in PSP brains are similar or different. However, if representation of 4R tau deposits is disease-specific, it is expected that molecular species of 4R tau itself is disease-specific as well. For example, it has been reported that asparagine at residue 279 of 4R tau is deamidated to aspartate in AD brains [13], while this posttranslational change is not robust in PSP/CBD brains [8]. Immunoprobes that may detect such AD-specific posttranslational changes may provide a straightforward strategy to demix PSP-type lesions and AD-type lesions in the same brain, in the same area or even within a single neuron at molecular level. If AD-type NFTs are characterized by paired helical filaments [29, 32] while PSP-type NFTs are characterized by straight fibrils [35] on electron microscopy, it will be exciting to examine how they are related to such biochemical differences, if any. Our hybrid approach to demix AD-type and PSP-type tau lesions may be corroborated by hybrid molecular demixing and electron microscopic studies, which will surely improve our mechanistic understanding of these diseases for more precise diagnosis and better management.
of pons; PU: putamen; RN: red nucleus; RPN: raphe nucleus; SN: substantia nigra; STN: subthalamic nucleus; TA: tuft-shaped astrocyte

Acknowledgements
Not applicable.

Funding
This work was supported by Grant-in-Aid for JSPS KAKENHI Grant Numbers 17H03555 (TU), 16K14572 (TU).

Authors' contribution
Study design: ME, YI, TU; Autopsy and neuropathological analyses: ME, YI, KH, TU; Immunohistochemical analyses and mapping: ME, MU, AN, SK, TU; Initial drafting: ME, SK; Final drafting: KH, SK, TU. All authors have read and approved the final manuscript.

Ethics approval and consent to participate
Written consent for autopsy and subsequent use of autopsied samples for research was obtained from the next of kin of each patient. The ethics committee of Tokyo Metropolitan Institute of Medical Science approved this study. (#16–25).

Competing interests
The authors declare that they have no interests.

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Received: 11 February 2019 Accepted: 22 March 2019
Published online: 06 May 2019

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