Double staining of elastic fibre and immunohistochemistry is helpful to differentiate pleural invasion of lung cancer

Visceral pleural invasion (VPI) is one of the critical determining factors for pathological stage of lung cancer. The pleural invasion (PL) status can be divided into four categories: PL0, PL1, PL2 and PL3, depending on the spatial relationship between the tumour tissue and outer elastic layer of pleura, visceral pleura surface and parietal pleura, respectively. Elastic stains are recommended if it is difficult to distinguish PL status by H&E staining. As reported, 19 of 100 lung cancer specimens were redefined as PL1 which were formerly deemed as PL0 by HE stain only, changing the stage from T1 (stage IA) to T2 (stage IB), which is essential for chemotherapy after surgery. Nonetheless, even assisted by elastic stains, VPI assessment is still difficult in some scenarios marked by multiple layers of elastic fibres, inflammatory infiltrates, fibrotic proliferation or carbon deposition which makes the location of scattered tumour cells barely discernible. To solve this technical problem, we developed a double staining method to show the spatial relationship between tumour cells and elastic fibres. Specifically speaking, the elastic fibres and tumour cells are stained and displayed by different colours on one slide, to determine whether the pleura is invaded.

Double staining process: Formalin-Fixed, Paraffin-Embedded (FFPE) blocks were sectioned by thickness 3–5 μm for double staining. Step one is routine immunohistochemistry staining, including: (1) antigen retrieval for 23 min in 97°C, (2) washing with distilled water, (3) 3% hydrogen peroxide solution incubated for 10 min to remove endogenous peroxidase, (4) rinsing with phosphate balanced solution (PBS) 3 min for three times, (5) add primary antibody (thyroid transcription factor-1 (TTF-1), and PanCK (AE1/AE3), (Zhongshan, Beijing, China)) and incubate at room temperature for 40–60 min, (6) rinsing with PBS, add the secondary antibody (Horseradish Peroxidase (HRP)) and incubate for 10–15 min at room temperature, (7) rinsing with PBS 3 min for three times and then diaminobenzidine (DAB) colour development. Second step is double dyeing of elastic fibre using a Fuxi Assay (Fuxi Bio, Guiyang, China) according to the instruction for 10–15 min, then 70% ethanol is used for counterstaining with haematoxylin. Finally, dehydrate and transparently seal after washing.

TTF-1 or AE1/AE3 shows a dark brown immunohistochemistry (IHC) signal, with the former exclusively nuclear stained and the latter localised to the cytoplasm. Elastic fibres are stained in purple while all the nuclei are stained blue with haematoxylin, including tumour cells, stromal cells, lymphocytes and so on. Figure 1 shows a nodule scanned in CT with slight pleural stretch, which is hard to discern in routine H&E slides since the elastic fibres are intact stained in an ambiguous pink architecture, but with the help of double staining, we can easily recognise tumour cells attached on the surface of pleura for a correct diagnosis for PL2. Figure 2 shows another difficult case for VPI status. (A) H&E staining shows the pleura full of fibres and capillaries (red arrow) with an acinar structure protruding towards the surface of pleura (thin black arrow, ×100). Double staining of AE1/AE3 (B, dark brown in cytoplasm), TTF-1 (C, dark brown in nucleus) and elastic fibre (in purple, ×100) confirms the protruding acinar structure to be adenocarcinoma (thin dark arrow from A to C) and further confirmed a capillary-like adenocarcinoma area (bold dark arrow from A to C) so as to achieve a correct diagnosis of PL1. PL, pleural invasion; TTF-1, thyroid transcription factor-1; VPI, visceral pleural invasion.
a more difficult case, from the H&E slide, we can hardly recognise elastic fibre and cancer cells from the conspicuous hyperplastic fibres and capillaries. However, our double staining method can easily identify the location of adenocarcinoma cells and elastic fibres and screen out the proliferative capillaries, which are obviously similar to acinar adenocarcinoma on H&E section.

Compared with the traditional elastic fibre staining, our double staining method has the following advantages: (1) easy interpretation, high repeatability and less controversy. (2) Working time is shorter for 2/3 than that of immunohistochemical staining and elastic fibre staining alone (2.5 hours vs 6 hours). (3) The cost is small. Our method is worth popularising for replacing routine elastic fibre staining to make a more correct diagnosis for pleural invasion.

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