Abstract

Background: Telepathology can potentially be utilized as an alternative to having on-site pathology services for rural and regional hospitals. The goal of the study was to validate two small-footprint desktop telepathology systems for remote parathyroid frozen sections.

Subjects and Methods: Three pathologists retrospectively diagnosed 76 parathyroidectomy frozen sections of 52 patients from three pathology services in Australia using the “live-view mode” of MikroScan D2 and Aperio LV1 and in-house direct microscopy. The final paraffin section diagnosis served as the “gold standard” for accuracy evaluation. Concordance rates of the telepathology systems with direct microscopy, inter-pathologist and intra-pathologist agreement, and the time taken to report each slide were analyzed. Results: Both telepathology systems showed high diagnostic accuracy (>99%) and high concordance (>99%) with direct microscopy. High inter-pathologist agreement for telepathology systems was demonstrated by overall kappa values of 0.92 for Aperio LV1 and 0.85 for MikroScan D2. High kappa values (from 0.85 to 1) for intra-pathologist agreement within the three systems were also observed. The time taken per slide by Aperio LV1 and MicroScan D2 within three pathologists was about 3.0 times (P < 0.001, 95% confidence interval [CI]: 2.8–3.2) and 7.7 times (P < 0.001, 95% CI: 7.1–8.3) as long as direct microscopy, respectively, while MikroScan D2 took about 2.6 times as long as Aperio LV1 (P < 0.001, 95% CI: 2.4–2.7). All pathologists evaluated Aperio LV1 as being more user-friendly.

Conclusions: Telepathology diagnosis of parathyroidectomy frozen sections through small-footprint desktop systems is accurate, reliable, and comparable with in-house direct microscopy. Telepathology systems take longer than direct microscopy; however, the time taken is within clinically acceptable limits. Aperio LV1 takes shorter time than MikroScan D2 and is more user-friendly.

Keywords: Frozen sections, parathyroidectomy, telepathology, time taken, validation study

INTRODUCTION

Access to the medical services continues to be an issue for many rural and regional areas in countries such as Australia. In the Australian context, when frozen sections are provided to the hospitals without an in-house laboratory, a pathologist has to travel to the operating theatre, setup the equipment, and wait for and then process the specimen. The time taken by the pathologist in diagnosing the prepared slide(s) is a fraction of the total time expended in any particular case. The utilization of new and current technology to bridge this gap in a cost-effective and safe manner is of great interest to public and private health services. Telepathology has the ability to minimize the nonprofessional “down-time” and provide intraoperative consultations to the hospitals that do not have an in-house pathologist and may prove to be cost-effective.

In the 1980’s, telepathology was first used to deliver pathology services remotely,[1] and since then there have been multiple technological advances.[2] Till date, a number of modalities, such as sending of selected static images, videotelemicroscopy, robotic microscopy, whole-slide imaging (WSI), and hybrid...
devices with both WSI and robotic microscopy capacities, have been used in telepathology for consultation, diagnosis, education, and quality assurance.\(^3\) Recently developed are hybrid desktop systems which are compact, cost-effective, and provide remotely controlled live imaging.

Validation studies using various telepathology systems for the primary diagnosis of paraffin and frozen sections of various tissue types have been reported, and we were only able to identify two reports which included parathyroid cases (≤5 cases) in among frozen sections of other organs.\(^{3,4,5}\) A very recent large sample-sized study of WSI of paraffin-embedded rather than frozen section specimens included more parathyroid cases (10 cases) and demonstrated its noninferiority to conventional microscopy for primary diagnosis.\(^6\) However, no study has specifically addressed the use of telepathology utilizing desktop-sized hybrid devices for parathyroid frozen sections.

International guidelines recommend validating a telepathology system for its intended use before formally using it to deliver a service.\(^7,8\) The purpose of this study was to validate two commercially available desktop telepathology systems for remote parathyroidectomy frozen section interpretation.

**Subjects and Methods**

**Ethics and case series**

Ethics committee approval was obtained from the Western Sydney Local Health District Human Research Ethics Committee (HREC Ref: (4275) AU RED LNR/15/WMEAD/153).

**Three pathology services in Sydney**. Austpath Laboratories (Austpath), Sydney South West Pathology Service (SSWPS), and Institute for Clinical Pathology and Medical Research (ICPMR), each submitted 20 consecutive parathyroidectomy cases. The participating pathologists were not involved in the selection of the cases. At Austpath, the frozen section specimens were accessioned from June 1, 2007 to April 30, 2015, at SSWPS from February 1, 2015 to March 31, 2016, and at ICPMR from April 1, 2014 to April 30, 2015. No additional inclusion or exclusion criteria were applied. This resulted in a total of 60 parathyroidectomy cases and 86 frozen sections (25, 37, and 24 from each service, respectively). All cases were de-identified and assigned research accession numbers. The final paraffin section diagnosis for each frozen section specimen was recorded from the archived report. According to the paraffin section diagnoses, the final sample consists of 36 parathyroid adenomas, 26 parathyroid hyperplasias, 18 normal parathyroids, two lymph nodes, two normal thyroid tissues, one fat tissue, and one with unclear recording of paraffin section diagnosis.

**Telepathology systems**

Two compact, desktop sized, low-volume white light whole-slide scanners with both WSI and live-view modes were installed in Austpath as the central facility. MikroScan D2 (11 × 13 × 8 inches in size with two-slide loading capacity, MikroScan Technologies, Vista, California, USA) was initially installed in Austpath in 2013 for evaluation in a van-based mobile frozen section service. Aperio LV1 (17.9 × 15.9 × 20.5 inches in size with four-slide loading capacity, Leica Biosystems, Vista, California, USA) was loaned for 10 days in Austpath to perform this study in 2016. Each scanner included a motorized microscope with a set of objective lenses. Although WSI had been successfully demonstrated for frozen section interpretation by others,\(^{12-15}\) we adopted their live-view mode instead of WSI during the study. WSI requires slide scanning and transfer of large files, which were more time-consuming in these two low-volume telepathology systems in addition to the challenge of internet congestion in the area where the study was conducted (4G connection was used). The live-view mode enabled remote users to navigate the slide and control the focus. This can be more useful when evaluating frozen sections which, not uncommonly, have tissue folds\(^7\) and are usually thicker than paraffin sections.

**Interpretation of parathyroidectomy frozen sections through telepathology and direct microscopy**

Three pathologists (Pathologist 1, Pathologist 2, and Pathologist 3 from Austpath, ICPMR and SSWPS, respectively) interpreted the parathyroidectomy frozen sections in the same order of slides using Aperio LV1, MikroScan D2, and in-house direct microscopy (direct microscopy) in sequence without knowledge of the previous frozen section and paraffin section diagnosis. No counterbalancing of the study design was applied. There were more than 30 years of board-certified anatomical pathology experience for Pathologists 1 and 3 and 6 years for Pathologist 2. Pathologist 1 specialized in head-and-neck pathology. The specialty areas of Pathologist 2 were skin, breast, soft tissue, and endocrine pathology. Pathologist 3 had special interest in endocrine, breast, gynecology, and gastrointestinal pathology. All the three pathologists had no previous experience in digital pathology reporting except for reading WSI cases for a quality assurance program in which virtual microscope images have been used as survey material since 2006. A short machine training session of each telepathology system had been given to each pathologist before the formal conduction of the study. As per the guideline,\(^7\) a washout period of ≥2 weeks between each examination was observed.

During each examination, one trained histotechnologist assisted with setting up the local scanner and computer, connecting to the remote computer through a third-party web conferencing application (WebEx Productivity Tools, [Cisco System Inc., USA] for MikroScan D2 and TeamViewer 11 [Team Viewer GmbH, Göppingen, Germany] for Aperio LV1), and loading the slides (one slide at a time). The stopwatch was started at the moment when the slide was loaded. The pathologist then remotely controlled the robotic microscope and viewed images on his personal computer (Pathologist 1: Same building different room; Pathologist 2: 1 kilometer away; and Pathologist 3: 20 kilometers away). Once the diagnosis was made, the pathologist typed the diagnosis into the web conferencing module. The histotechnologist then stopped the
stopwatch and recorded the diagnosis and the time taken to reach the diagnosis.

For direct microscopy, each pathologist interpreted the slides through a conventional microscope in his own office. The on-site histotechnologist started the stopwatch as soon as the pathologist received the slide and stopped when the pathologist communicated the diagnosis.

**Data analysis**

The final paraffin section diagnosis for each slide was standardized as either “parathyroid” (including parathyroid adenoma, hyperplasia, and normal parathyroid) or “not parathyroid” (including lymph node, thyroid, and fat tissue) and served as the reference (“gold standard”) to determine the diagnostic accuracy. The reviewed frozen section diagnoses were similarly classified into the above two categories with the addition of “defer to paraffin section.”

Microsoft Excel 2010 (Microsoft Corporation) and IBM SPSS Statistics version 24.0 (IBM Corporation, Armonk, New York, USA) were used to analyze the data. The validity of all three systems and the concordance rates of telepathology systems with direct microscopy were calculated. The inter-pathologist and intra-pathologist agreement was evaluated by overall kappa values (Fleiss’ kappa\(^6\)) for three and Cohen’s kappa\(^{17}\) for two raters. Logarithmic transformation (In) of time taken before analysis was applied to stabilize the variance of data. Repeated measures analysis of variance (ANOVA) of transformed data was conducted with “system” treated as a three-level factor within the three-level “pathologist” factor. Where a significant within “slides” effect was observed, the Bonferroni-corrected pairwise comparisons were performed.  

**RESULTS**

**Data summary**

Eighty-six frozen sections were accessioned and of these ten were excluded for technical reasons as follows: one for incorrect accession (not frozen section), four for failure of loading onto these telepathology systems because of broken and repaired slides, one for unclear record of final diagnosis, and four for significantly faded stains. Seventy-six frozen sections (72 “parathyroid” and 4 “not parathyroid” according to the final paraffin diagnoses) from 52 cases were analyzed.

**Validity of telepathology systems**

As shown in Tables 1 and 2, all three systems showed very high diagnostic accuracy in identifying parathyroid from frozen section specimens (100.0% for direct microscopy, 99.6% for Aperio LV1, and 99.1% for MikroScan D2). One normal parathyroid specimen (FST33B) was deferred by one pathologist in Aperio LV1, and there were no false diagnoses and the 100.0% of positive predictive value and negative predictive value (NPV) confirmed the efficiency of this system. Two normal parathyroid specimens (FST33B and FST19D) were falsely diagnosed as either “no parathyroid tissue” or “thymus” by one pathologist through MikroScan D2, which led to a false-negative rate (FNR) of 0.9% and NPV of 85.7% for this system. Retrospective analysis revealed that FST33B was mainly fat, and the pathologist had commented that it was a “terrible section.” FST19D was falsely diagnosed as “not parathyroid, thymus” by one pathologist through MikroScan D2 but correctly diagnosed as “parathyroid” by the same pathologist through the other two systems.

**Reliability of telepathology systems**

When compared with direct microscopy, an overall concordance rate of 99.6% (1/228 discrepancy due to a deferral through telepathology) for Aperio LV1 and 99.1% for MikroScan D2 (2/228 discrepancies due to false-negative diagnosis) was observed. As shown in Table 3, the direct microscopy showed a perfect inter-pathologist agreement with an overall kappa value of 1. Aperio LV1 and MikroScan D2 also presented high inter-pathologist concordance with an overall kappa value of 0.92 and 0.85 (both >0.75), respectively.

The overall kappa values of intra-pathologist agreement [Table 3] also demonstrated a high replicability of frozen section diagnosis of parathyroidectomy specimens for each pathologist through three different systems (0.85–1, all >0.75).

**Turnaround times**

As shown in Figure 1 and Table 4, the median of time taken to reach the diagnosis of each parathyroid frozen section slide by pathologist 1, 2, and 3 was 45 s, 20 s, and 20.5 s through direct microscopy; 105 s, 60 s, and 78.5 s through Aperio LV1; and 185 s, 200 s, and 223 s through MikroScan D2, respectively. The statistical analysis of the log-transformed data, repeated measures ANOVA with Greenhouse-Geisser correction followed by Bonferroni-corrected pairwise comparisons, revealed that Aperio LV1 and MikroScan D2 took about 3.0 times \((P < 0.001, 95\%\text{ CI}: 2.8–3.2)\) and 7.7 times \((P < 0.001, 95\%\text{ CI}: 7.1–8.3)\) as long as direct microscopy within three pathologists, respectively. MikroScan D2 took about 2.6 times as long as Aperio LV1 \((P < 0.001, 95\%\text{ CI}: 2.4–2.7)\).

**DISCUSSION**

In this study, we specifically validated two small-footprint, low-volume, hybrid telepathology systems for parathyroidectomy frozen section interpretation. We demonstrated a very high diagnostic accuracy of both systems (99.6% for Aperio LV1 and 99.1% for MikroScan D2...
Another pathologist (Pathologist 4) instead of Pathologist 2 evaluated all cases through Aperio LV1, as Pathologist 2 was on leaving during the time this system was accessed. PPV: Positive predictive value, NPV: Negative predictive value, FPR: False-positive rate, FNR: False-negative rate

Comparison of diagnostic accuracy between telepathology and direct microscopy

Parathyroidectomy frozen sections can be less challenging with a narrower range of diagnostic possibilities than those in neuropathology and general surgery. In line with our usual diagnostic practice, the frozen section diagnosis in this study was categorized as “parathyroid,” “not parathyroid,” or “defer to paraffin section.” Two normal parathyroid specimens were incorrectly diagnosed as “not parathyroid” through MikroScan D2, which led to an FNR of 0.9%. One of these could be attributed to the sample and section quality, as it was recorded that the specimen was fatty and the section was obtained with difficulty. The other was a diagnostic error of the pathologist that the specimen was fatty and the section was obtained with difficulty. The other was a diagnostic error of the pathologist rather than an inferior performance of the telepathology system.

When compared to the direct microscopy, an overall concordance rate of 99.6% for Aperio LV1 and 99.1% for MikroScan D2 was recorded. The overall kappa values of intra-pathologist concordance within these two telepathology systems and direct microscopy reached 0.85–1 for each pathologist. The inter-pathologist agreement for these two telepathology systems similarly reached a high level (overall kappa value of 0.92 for Aperio LV1 and 0.85 for MikroScan D2). This means that both telepathology systems were comparable to the direct microscopy and were sufficiently reliable to be used by different pathologists or the same pathologist at different times. Similar high levels of concordance between dynamic/WSI telepathology and in-house conventional microscopy in frozen section diagnosis have also been reported.¹

In this study, the diagnosis by both telepathology systems took longer than direct microscopy. Aperio LV1 took 3.0 times, and MikroScan D2 took 7.7 times as long as direct microscopy (both P < 0.001). Preliminary scanning, transmission of the remote operation commands and response of the robotic microscope to the commands, digitizing and transmission of the images, and prolongation of transmission due to internet congestion contributed to the longer time taken by the telepathology systems.

Published turnaround time for the frozen section by telepathology varies and depends mainly on

### Table 2: Validity of direct microscopy, Aperio LV1, and MikroScan D2 (%, n = 228)

| System         | Defer | Sensitivity | Specificity | PPV  | NPV  | FPR  | FNR  | Accuracy |
|----------------|-------|-------------|-------------|------|------|------|------|----------|
| Direct microscopy | 0.0   | 100.0       | 100.0       | 100.0| 100.0| 0.0  | 0.0  | 100.0    |
| Aperio LV1†    | 0.4   | 99.5        | 100.0       | 100.0| 100.0| 0.0  | 0.0  | 99.6     |
| MikroScan D2   | 0.0   | 99.1        | 100.0       | 100.0| 85.7 | 0.0  | 0.9  | 99.1     |

†Another pathologist (Pathologist 4) instead of Pathologist 2 evaluated all cases through Aperio LV1, as Pathologist 2 was on leaving during the time this system was accessed.

### Table 3: Inter-pathologist and intra-pathologist agreement of two telepathology systems and direct microscopy (Fleiss’ kappa)

| Agreement                | n   | Overall kappa | 95% CI          |
|--------------------------|-----|---------------|-----------------|
| Inter-pathologist        |     |               |                 |
| Direct microscopy        | 76  | 1             | 0.87-1.13       |
| Aperio LV1               | 76  | 0.92          | 0.80-1.04       |
| MikroScan D2             | 76  | 0.85          | 0.71-0.98       |
| Intra-pathologist        |     |               |                 |
| Pathologist 1            | 76  | 1             | 0.87-1.13       |
| Pathologist 2†           | 76  | 1             | 1-1             |
| Pathologist 3            | 76  | 0.85          | 0.72-0.98       |

†The overall kappa value was calculated according to Cohen’s kappa, as Pathologist 2 only participated in the evaluation through two systems (direct microscopy and MikroScan D2). CI: Confidence interval.

Figure 1: Boxplot of time taken per slide of parathyroid frozen sections reviewed by three pathologists through three systems (n = 76). As the data showed large variance in some groups, a logarithmic transformation of the data was applied before the repeated measures analysis of variance and post hoc tests. The dark line in the middle of the boxes is the median of the values of time taken. The small circles (○) represent the outliers with values more than 1.5× interquartile range. The asterisks (⋆) are extreme outliers with values more than 3× interquartile range. ³Another pathologist (Pathologist 4) instead of Pathologist 2 interpreted all the 76 slides through Aperio LV1.) s, seconds; PG, Pathologist; IQR: interquartile range.

The overall kappa value was calculated according to Cohen’s kappa, as Pathologist 2 only participated in the evaluation through two systems (direct microscopy and MikroScan D2). CI: Confidence interval.

The overall kappa value was calculated according to Cohen’s kappa, as Pathologist 2 only participated in the evaluation through two systems (direct microscopy and MikroScan D2). CI: Confidence interval.
the specimens interpreted, devices used, and pathologists’ experience in frozen section diagnosis and using digital images. The time taken by telepathology diagnosis of parathyroid frozen sections in this study (median value around 4 min per slide) was shorter than those reported in neuropathology (around 10 min per slide[10,26]) and similar to those recorded in studies of mixed tissue types (around 3 min per slide[15]).

Evans et al.[10] recorded a significantly lower time taken by their newly implemented WSI scanner than the previous robotic microscopy system during their routine telepathology frozen section practice. In this study, we reviewed the same cases through two similar telepathology systems and still observed a significantly shorter time taken for Aperio LV1 than for MikroScan D2 (P < 0.001). A more user-friendly operating experience with Aperio LV1 was also informally recorded (no questionnaires were performed). MikroScan D2 and Aperio LV1 were released in 2010 and 2015, respectively.[27,28] The 5-year gap in an actively evolving field could be one of the main reasons for the differences.

Besides the issue of the longer time taken, a broken, oversized, or suboptimal slide can be a challenge to the telepathology systems. However, in the clinical situation, this will be rectified by cutting and staining another section.

Our study has demonstrated the accuracy and reliability of telepathology in the frozen section diagnosis of parathyroidectomy specimens. For the surgeons using this service, a 6-min prolongation was acceptable and did not adversely impair best patient care. Given this short delay, regional and country hospitals would be able to consider this as a financially viable alternative to having an on-site frozen section service. However, as the order effects of the repeated measures design of the study was not controlled by counterbalancing, unplanned change of the participating pathologist occurred for part of the study, and relatively few “not parathyroid” cases were included as subjects in the sample, systematic and sampling biases might influence the outcomes of this study. Additional studies with more powerful design and greater case numbers and evaluation of costs will be needed to confirm and reinforce our findings before implementing telepathology for parathyroid frozen sections more widely in clinical practice. Training on any specific telepathology system is still necessary, and each laboratory will need to formulate its own protocols.

**Conclusions**

Telepathology diagnosis of parathyroidectomy frozen sections through small-footprint desktop systems is accurate, reliable, and comparable with in-house direct microscopy. Telepathology systems take longer than direct microscopy; however, the time taken is within clinically acceptable limits. Aperio LV1 takes shorter time than MikroScan D2 and is more user-friendly.

**Acknowledgments**

Dr. Mahatab Farzin, MD, FRCPA. Department of Anatomical Pathology, NSW Health Pathology, Orange Campus, evaluated all cases using the Aperio LV1 and her contribution to the study is acknowledged.

**Financial support and sponsorship**
Nil.

**Conflicts of interest**
There are no conflicts of interest.

**References**

1. Pradhan D, Monaco SE, Parwani AV, Ahmed I, Duboy J, Pantanowitz L, et al. Evaluation of panoramic digital images using Panoptiq for frozen section diagnosis. J Pathol Inform 2016;7:26.
2. Pantanowitz L, Wiley CA, Demetris A, Lesniak A, Ahmed I, Cable W, et al. Experience with multimodality telepathology at the university of Pittsburgh medical center. J Pathol Inform 2012;3:45.
3. Evans AJ, Salama ME, Henricks WH, Pantanowitz L. Implementation of whole slide imaging for clinical purposes: Issues to consider from the perspective of early adopters. Arch Pathol Lab Med 2017;141:944-59.
4. Bauer TW, Slaw RJ, McKenney JK, Patil DT. Validation of whole slide imaging for frozen section diagnosis in surgical pathology.
J Pathol Inform 2018, 1:41

J Pathol Inform 2015;6:49.
5. Kaplan KJ, Burgess JR, Sandberg GD, Myers CP, Bigott TR, Greenspan RB, et al. Use of robotic telepathology for frozen-section diagnosis: A retrospective trial of a telepathology system for intraoperative consultation. Mod Pathol 2002;15:1197-204.

6. Mukhopadhyay S, Feldman MD, Abels E, Ashfaq R, Beltaifa S, Cacciabeve NG, et al. Whole slide imaging versus microscopy for primary diagnosis in surgical pathology: A Multicenter blinded randomized noninferiority study of 1992 cases (Pivotal study). Am J Surg Pathol 2018;42:39-52.

7. Canadian Association of Pathologists Telepathology Guidelines Committee; Bernard C, Chandrananth SA, Cornell IS, Dalton J, Evans A, et al. Guidelines from the Canadian association of pathologists for establishing a telepathology service for anatomic pathology using whole-slide imaging. J Pathol Inform 2014;5:15.

8. Pantanowitz L, Dickinson K, Evans AJ, Hassell LA, Henricks WH, Lennerz JK, et al. American telemedicine association clinical guidelines for telepathology. J Pathol Inform 2014;5:39.

9. Têtu B, Perron E, Louahlia S, Paré G, Trudel MC, Meyer J, et al. The Eastern Québec telepathology network: A three-year experience of clinical diagnostic services. Diagn Pathol 2014;9 Suppl 1:S1.

10. Evans AJ, Chetty R, Clarke BA, Croul S, Ghazarian DM, Kiehl TR, et al. Primary frozen section diagnosis by robotic microscopy and virtual slide telepathology: The university health network experience. Hum Pathol 2009;40:1070-81.

11. Fallon MA, Wilbur DC, Prasad M. Ovarian frozen section diagnosis: Use of whole-slide imaging shows excellent correlation between virtual slide and original interpretations in a large series of cases. Arch Pathol Lab Med 2010;134:1020-3.

12. Ramey J, Fung KM, Hassell LA. Use of mobile high-resolution device for remote frozen section evaluation of whole slide images. J Pathol Inform 2011;2:41.

13. Gould PV, Saikali S. A comparison of digitized frozen section and smear preparations for intraoperative neurotelepathology. Anal Cell Pathol (Amst) 2012;35:85-91.

14. Perron E, Louahlia S, Nadeau L, Boilard F, Ing M, Orain M, et al. Telepathology for intraoperative consultations and expert opinions: The experience of the Eastern Québec telepathology network. Arch Pathol Lab Med 2014;138:1223-8.

15. Elhassan ME, Anikin V, Beddow E, McGonigle N, Kyparissopoulos D, Rice A, et al. Assessment of intraoperative frozen sections diagnosed based on remote reporting by telepathology. Pathology 2014;46 Suppl 2:S63.

16. Zaiontz C. Fleiss’ Kappa. Real Statistics Using Excel; Posted 02 November, 2013. Available from: http://www.real-statistics.com/reliability/fleiss-kappa/. [Last accessed on 2017 Feb 28].

17. Zaiontz C. Cohen’s Kappa. Real Statistics Using Excel; Posted 26 December, 2012; Available from: http://www.real-statistics.com/reliability/cohens-kappa/. [Last accessed on 2017 Feb 28].

18. Wellnitz U, Binder B, Fritz P, Friedel G, Schwarzmann P. Reliability of telepathology for frozen section service. Anal Cell Pathol 2000;21:213-22.

19. Wiley CA, Murdoch G, Parwani A, Cudahy T, Wilson D, Payner T, et al. Interinstitutional and interstate teleneuropathology. J Pathol Inform 2011;2:21.

20. Horbinski C, Wiley CA. Comparison of telepathology systems in neuropathological intraoperative consultations. Neuropathology 2009;29:655-63.

21. Vitkovski T, Bluiya T, Esposito M. Utility of telepathology as a consultation tool between an off-site surgical pathology suite and affiliated hospitals in the frozen section diagnosis of lung neoplasms. J Pathol Inform 2015;6:55.

22. Ribback S, Flessa S, Gromoll-Bergmann K, Evert M, Dombrowski F. Virtual slide telepathology with scanner systems for intraoperative frozen-section consultation. Pathol Res Pract 2014;210:377-82.

23. Gifford AJ, Colebatch AJ, Litkouhi S, Hersch F, Warzeca W, Snook K, et al. Remote frozen section examination of breast sentinel lymph nodes by telepathology. ANZ J Surg 2012;82:803-8.

24. Słodkowska J, Pankowski J, Siemiatkowska K, Chyczewski L. Use of the virtual slide and the dynamic real-time telepathology systems for a consultation and the frozen section intra-operative diagnosis in thoracic/pulmonary pathology. Folia Histochim Cytobiol 2009;47:679-84.

25. Hitchcock CL, Hitchcock LE. Three years of experience with routine use of telepathology in assessment of excisional and aspirate biopsies of breast lesions. Croat Med J 2005;46:449-57.

26. Hutarew G, Schlicker HU, Idriceanu C, Strasser F, Dietze O. Four years experience with teleneuropathology. J Telemed Telecare 2006;12:387-91.

27. Kaplan KJ. Another Scanner Company?; Posted 03 November, 2010. Available from: http://www.tissuepathology.com/2010/11/03/my-entry-5/#axzz4xdllyzoE. [Last accessed on 2017 Nov 02].

28. Leica Biosystems. Leica Biosystems Introduces the Aperio LV1 – Live View, Remote, Digital Pathology System for Research Use Only. Leica Biosystems; 01 October, 2015. Available from: http://www.leicabiosystems.com/news-events/news-details/article/leica-biosystems-introduces-the-aperio-lv1-live-view-remote-digital-pathology-system-for-research/News/detail/. [Last accessed on 2017 Nov 02].