RESEARCH ARTICLE

Errors in Surgical Pathology Reports: a Study from a Major Center in Pakistan

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Abstract

**Background:** Errors in surgical pathology diagnosis can have serious consequences for the patient. Since the final product of a surgical pathology lab is the report, errors can be picked by reviewing reports of cases. **Aim:** To determine the frequency and types of error in surgical pathology reports of cases signed out in 2014 in a laboratory in Karachi, Pakistan. **Materials and Methods:** All surgical pathology reports in which changes were made in the original report after sign out and an amended report was issued were included. Errors included: (1) misinterpretations; (2) missing critical information; (3) erroneous critical information; (4) misidentification; and (5) typographic errors. **Results:** Errors were identified in 210 cases (0.37%). These comprised 199 formalin fixed specimens and 11 frozen sections. The latter represented 3.8% of a total of 2,170 frozen sections. Of the 11 frozen section errors, 10 were misinterpretations. Of the 199 permanent specimens, 99 (49.7%) were misinterpretations, 65 (32.7%) belonged to missing critical information category, 8 (4%) belonged to erroneous critical information category, 8 (4%) were misidentifications, 16 (8%) were typographic errors while 3 cases (1.5%) were other errors. Most misinterpretations occurred in the gastrointestinal, liver and pancreatic biliary tract (23.2%) and breast (13.1%). Another 87 cases were reviewed on the clinicians’ request. However diagnosis after review remained the same as the original diagnosis. In 49 out of these (56.3%), additional workup was performed at the time of the review. **Conclusions:** Our findings were similar to other published studies. We need to develop documented procedures for timely review of cases to detect errors.

**Keywords:** Surgical pathology - amended reports - errors - misinterpretations - omission of critical information

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Introduction

Errors in surgical pathology diagnosis can have serious consequences for the patient. Since the principal product of surgical pathology laboratory is the report, errors are mostly picked by reviewing reports of cases that have been dispatched. Errors are either picked by clinicians when the reports do not match the clinical diagnosis or by the pathologists themselves. The pathologists discover the error either by mandatory reviews of the slides and reports at the time of case referral to another institution for second opinion, or when reviewing the slides for a surgical pathology conference especially tumor boards or during regular audit of surgical pathology cases as part of a routine quality assurance exercise. The pathologists can also discover the error when reviewing the permanent sections of a case reported initially as a frozen section. Errors in surgical pathology can be major or clinically significant if they have adverse effects on the patient’s clinical management and minor or clinically insignificant if they do not have adverse consequences on the patient’s clinical management.

Errors in surgical pathology can be misinterpretations which include false positives or false negatives, errors in determining the biologic behavior of tumors whether benign or malignant or errors in determining the histogenesis of tumors. Errors can also be misidentifications of the patient, tissue, specimen laterality and/or errors in identification of anatomic localization of the specimen. Errors in surgical pathology reports also include the omission of critical information or inclusion of wrong information in the reports. Another common type of error in reports are typographical errors such as those of spellings, grammar, formatting etc.

All errors when discovered lead to a review of the case and any changes, corrections etc. are then issued as amended or addendum reports. In some cases, clinicians are not satisfied by the reports and request the pathologists to review the case. If on review, without or after additional workup, no error is detected and the original report stands, the fact is communicated in the form of an addendum report.

The aim of this study was to determine the frequency and types of errors in surgical pathology reports of cases signed out in 2014. The Section of Histopathology at Aga Khan University Hospital (AKUH) in Karachi is the
largest in Pakistan and we get specimens from all over the country. (Ahmad et al., 2014)

**Materials and Methods**

All surgical pathology reports signed out in 2014 in which changes were made in the original report to correct errors of any type after the case led been signed out and in which an amended report was subsequently issued, were included in the study. In addition, all those reports where after sign-out case was reviewed on the clinician’s request but the diagnosis after review remained unchanged (with or without additional workup) and in which an addendum report was issued to document the same, were also included in the study. Errors in surgical pathology reports were classified as follows:

1. Misinterpretations defined as inaccurate or incorrect diagnosis. These included those in which change in diagnosis occurred on the review of the case without performing any additional workup - “per se”; as well as those misinterpretations which occurred as a result of failure to perform appropriate immunohistochemical (IHC) and/or special stains, failure to submit and examine appropriate sections, failure to submit and examine adequate number of sections, failure to perform and examine deeper levels etc. at the time of the original sign out and which when performed at the time of review led to a significant change in diagnosis.

2. Missing critical information in reports defined as information the omission of which in surgical pathology reports was likely to have significant adverse impact on patient care. Examples include missing critical measurements such as distance of tumor from excision margins in malignant neoplasms, margin status i.e. positive or negative, missing tumor grade or stage, missing tumor size and other critical information for example omission of status of nodes or adenaxial involvement in endometrial carcinoma, omission of status of detrusor muscle involvement in urothelial carcinomas of urinary bladder, omission of information about extra prostatic extension in prostatic adenocarcinoma on radical prostatectomy specimens etc. Non neoplastic examples include omission of status of fungal stain in nasal polyps, omission of information regarding presence or absence of Helicobacter pylori in gastric mucosal biopsies etc.

3. Erroraneous critical information was defined as erroneous information the inclusion of which in surgical pathology reports was likely to have significant adverse on patient care. Examples include the incorrect tumor grade or stage, incorrect tumor size etc.

4. Misidentifications defined as errors in identifying a tissue correctly or errors made in determining the laterality of a specimen.

5. Typographic errors defined as spelling or grammatical errors which could lead to misinterpretation of surgical pathology reports. Computer formatting errors were also included in this category.

The data was reviewed by the two principal authors (ZA and RI). All data was recorded and analyzed using the SPSS 19.0 software package. Cytology specimens were not included in the study.

**Results**

A total of 57000 surgical pathology cases were signed out in 2014. A total of 297 reports were included in the study. Errors in surgical pathology reports were identified in 210 cases or 0.37%. These 210 cases included 11 cases in which frozen sections were requested and there were discrepancy between the original diagnosis rendered on frozen section and the final diagnosis given on permanent section. The total number of frozen section cases during the year was 2170 (out of 57000) or 3.8%. The discordance rate on frozen section cases was 0.51%. In addition to the 210 cases in which errors were identified, another 87 were reviewed on clinicians’ request. However, the diagnosis after review remained the same as the original diagnosis. In 49 out of these 87 cases (56.3%) additional workup was performed (for example deeper levels, additional sections, special stains, immunohistochemical stains etc. However, this additional workup did not result in a change in diagnosis. In 38 out of 87 cases (43.7%), cases were reviewed but no additional workup was done and there was no change in diagnosis as a result of the review. Out of 11 frozen section errors, 10 were interpretative errors or misinterpretations (Table 1) while in 1 case, a focus of carcinoma was seen in a sentinel node (from a patient with carcinoma breast) on deeper permanent level. A diagnosis of “negative for metastasis” had been rendered at the time of frozen section .All 11 frozen section errors were picked

| S.No. | Tissue/Organ | Frozen Section Diagnosis | Final Diagnosis (on permanent sections) |
|-------|--------------|--------------------------|----------------------------------------|
| 1     | Central Nervous System | Meningioma | Myxopapillary Ependymoma |
| 2     | Lymph Node | Reactive lymph node | Low grade Bcell lymphoproliferative disorder |
| 3     | Central Nervous System | High grade glioma | Metastatic carcinoma |
| 4     | Head & Neck. Tongue Squamous cell carcinoma: Margins | Margins: dysplastic | Margins: benign(no dysplasia seen) |
| 5     | Central Nervous System | low grade glioma | High grade glioma |
| 6     | Ovary | Benign serous cystadenoma | Borderline serous neoplasma* |
| 7     | Central Nervous System | Inflammation | Lymphoproliferative disorder |
| 8     | Central Nervous System | Lymphoproliferative disorder | High grade glioma |
| 9     | Head & Neck. Cheek Squamous cell carcinoma: Margins | Margins: dysplastic | Margins: benign(no dysplasia seen) |
| 10    | Peritoneal nodule | Atypical cells seen(signet ring cells) | Plasma cells(no atypical cells seen) |
Table 2. Errors Identified in Paraffin Embedded Specimen Reports. (n=199)

| S. No. | Error type                                                                 | Number | Percentage (%) |
|--------|------------------------------------------------------------------------------|--------|----------------|
| 1      | Misinterpretation due to failure to perform appropriate IHC stains          | 24     | 12.10%         |
| 2      | Misinterpretation due to failure to perform appropriate special stains      | 7      | 3.50%          |
| 3      | Misinterpretation due to failure to submit appropriate/adequate sections for histologic examination | 25     | 12.60%         |
| 4      | Misinterpretation due to failure to perform adequate deeper level/s         | 13     | 6.50%          |
| 5      | Misinterpretation per se i.e. not due to lack of any appropriate workup     | 30     | 15.10%         |
| 6      | Missing critical /important information in reports                          | 65     | 32.70%         |
| 7      | Erroneous critical /important information                                   | 8      | 4.00%          |
| 8      | Typographic errors                                                          | 16     | 8.00%          |
| 9      | Wrong identification/coding or laterality of tissue                         | 8      | 4.00%          |
| 10     | Wrong diagnostic term used                                                  | 1      | 0.50%          |
| 11     | Specimen not reported at all                                                | 2      | 1%             |

Table 3. Examples of Misinterpretations Occurring as a Result of Failure to Perform Appropriate IHC Stains

| S.no. | Original diagnosis                          | Corrected diagnosis                          | IHC stains performed on case review          |
|-------|---------------------------------------------|----------------------------------------------|---------------------------------------------|
| 1     | High grade Glioma                          | Metastatic poorly differentiated carcinoma   | CK AE1/AE3 positive, CK CAM 5.2 positive     |
| 2     | Diffuse large B Cell Lymphoma              | Blastoid variant of Mantle cell lymphoma     | CD20,LCA Positive, MUM1,CD56 Negative       |
| 3     | Plasma cell neoplasm                       | Diffuse large B Cell Lymphoma                | Calretinin,WT1,CK5/6 positive. TTF1 Negative |
| 4     | Poorly differentiated carcinoma            | Malignant Mesothelioma                       |                                             |
| 5     | Lung Adenocarcinoma                        | Colon carcinoma                              | TTF1:Negative, CK20:Positive                |

Table 4. Examples of Misinterpretations Occurring as a Result of Failure to Perform Appropriate Special Stains/Adequate Deeper Levels

| S. No. | Original diagnosis                                      | Corrected diagnosis                                      | Special stains performed on case review |
|--------|--------------------------------------------------------|----------------------------------------------------------|----------------------------------------|
| 1      | Nasal polyp (fungal stains not performed)               | Polyp along with septate fungal hyphae                   | PAS/PASD Positive                       |
| 2      | Nasal polyp (fungal stains not performed)               | Polyp along with septate fungal hyphae                   | PAS/PASD Positive                       |
| 3      | Reactive gliosis brain (fungal stains not done)         | Septate fungal hyphae seen                               | PAS/PASD Positive                       |
| 4      | Cecal biopsy No granuloma on initial slide (level not done) | Granuloma seen on deeper levels. Possibility of Tuberculosis raised. | Deeper level was performed on case review. |
| 5      | Rectal polyp. No adenomatous change on initial section(no level performed) | Adenomatous change was seen on deeper levels. Tubular Adenoma | Deeper level was performed on case review. |
| 6      | Gastric biopsy from mass in stomach No atypical cells seen | Atypical cells seen on deeper levels. Diagnosis of carcinoma was given ganglion cells seen on multiple deeper levels | Deeper levels and Mucin stain were performed on case review. |
| 7      | Colon biopsy for ganglion cells No ganglion cells seen on initial two levels. | Tumor invades through cortical bone. | Multiple deeper levels performed on case review. |

Table 5. Examples of Misinterpretations Occurring as a Result of Failure to Submit Appropriate/Adequate Sections for Microscopic Examination

| S. No. | Original Diagnosis                                      | Corrected Diagnosis                                      | Additional sections submitted at time of review |
|--------|--------------------------------------------------------|----------------------------------------------------------|-----------------------------------------------|
| 1      | Breast Carcinoma T2N0                                   | Breast Carcinoma T2N1                                     | Missed at time of initial gross, nodes positive |
| 2      | Gall bladder, Chronic cholecystitis                      | Adenoma with low grade dysplasia                         | Polyp missed at the time of initial gross     |
| 3      | Ovarian Serous cystadenoma                              | Borderline serous tumor                                  | Additional sections revealed borderline areas, but no stromal invasion seen. |
| 4      | Breast specimen: lesion missed at initial gross          | Carcinoma ,T1N0                                           | Lesion seen and section submitted at time of review. |
| 5      | Urinary Bladder Carcinoma, Urothelial high grade;Detrussor muscle not identified in the original multiple sections | Invasive papillary urothelial carcinoma                   | Additional sections reveal detrusor muscle involved by the tumor |
| 6      | Breast carcinoma, DCIS.No invasion seen in multiple sections submitted. | Invasive Ductal carcinoma, NOS | Additional sections reveal areas of invasion |
| 7      | Carcinoma oral cavity; sections from bone not submitted | Tumor invades through cortical bone                        | Sections from bone were submitted and show involvement by tumor. |
Table 6. Examples of Misinterpretations Occurring Per Se

| S. No. | Original Diagnosis | Corrected Diagnosis(without any additional workup) |
|--------|-------------------|--------------------------------------------------|
| 1      | Pancreas; Mixed ductal and endocrine tumor | Chronic Pancreatitis |
| 2      | Breast; invasive lobular carcinoma on core biopsy | invasive ductal carcinoma on mastectomy specimen |
| 3      | Colon endoscopic biopsy: Non specific colitis | Inflammatory bowel Disease(ulcerative colitis) |
| 4      | Thigh muscle; Neurofibroma | Spindle cell lipoma |
| 5      | Bronchial biopsy: Tuberculosis | Sarcoidosis |
| 6      | Spinal meninges: Arachnoid cyst | Enterogenous cyst(Neuroenteric cyst) |
| 7      | Bone: fracture changes | Benign bone cyst |
| 8      | Gastrointestinal tract: Nonspecific colitis | Amyloidosis |
| 9      | Brain: Oligodendroglia,WHO grade 2 | Anaplastic Oligodendroglia,WHO grade 3 |
| 10     | Duodenum: mild nonspecific duodenitis | Brunner’s gland hyperplasia |
| 11     | Skin biopsy: Atopic dermatitis | Neonatal SLE(systemic lupus erythematosus) |
| 12     | Endometrial curettings: Exogenous hormonal effect | Excessive estrogen effect (Exogenous hormones not given) |

Table 7. Distribution of Misinterpretations in Various Organs (n=99)

| S. No. | Organ/System | Number | Percentage (%) |
|--------|--------------|--------|----------------|
| 1      | Gastrointestinal tract(including liver,pancreas & biliary tract) | 23     | 23.20% |
| 2      | Breast       | 13     | 13.10% |
| 3      | Lungs,pleura,mediastinum | 10     | 10.10% |
| 4      | Lymph nodes  | 9      | 9.10%  |
| 5      | Head & Neck (including nasal, paranasal sinuses, salivary gland, oral cavity, gums etc.) | 9      | 9.10%  |
| 6      | Kidney &urinary bladder | 8      | 8.10%  |
| 7      | Female genital tract | 8      | 8.10%  |
| 8      | Bone and Soft tissue | 7      | 7.10%  |
| 9      | Brain        | 6      | 6.10%  |
| 10     | Skin         | 4      | 4.00%  |
| 11     | Male Genital Tract | 2      | 2.00%  |

by the pathologist when reviewing the permanent sections.

The remaining 199 cases in which errors occurred were all paraffin embedded specimens. The large majority of these, 180 cases out of 199(90.5%), were detected when clinicians requested a review. Only 19 (9.5%) were detected by the pathologist who reported the case. Borderline features were seen in the permanent section of the original section submitted at the time of frozen section as well as in the additional sections submitted subsequently. However, in these cases no evidence of invasion was seen in any of the sections provided by the Pathologist.

Out of 199 cases, 99 (49.7%) errors occurred as a result of misinterpretations. These included errors in which misinterpretations occurred as a result of failure to perform appropriate immunohistochemical (IHC) stains and /or special stains, failure to submit and examine adequate number of sections, failure to perform deeper levels etc., all of which led to major changes in diagnosis. In addition, a number of cases were misinterpreted per se that is review of the cases on the clinicians’ request led to a change in diagnosis based on the workup originally performed without the need for performing any additional workup. In 65 cases (32.7%), critical /important information was missing in the reports which was later added. Another important group was that of typographical errors which accounted for 16 cases (8%).The details of these errors are shown in Table 2.

Examples of misinterpretations occurring as a result of failure to perform appropriate IHC stains are shown in Table 3. Examples of misinterpretations occurring as a result of failure to submit appropriate /adequate sections for microscopic examination are shown in Table 4. Examples of misinterpretations occurring per se are shown in Table 6. Distribution of interpretation errors (misinterpretations) in various organs of the body is given in Table 7.

Discussion

Recently, the College of American Pathologists and the Association of Directors of Anatomic and Surgical Pathology formed an expert panel which was asked to formulate recommendations based on existing evidence that additional case reviews of surgical pathology and cytology cases detect diagnostic errors. The panel drafted five recommendations for which there was strong agreement between the members. The first of these recommendations was that anatomic pathologists should develop documented procedures for timely review of selected cases to detect errors and should monitor and document the results of these reviews, and in case serious errors are detected, take steps to reduce the same.(Nakhleh et al., 2015).

Zarbo et al. (2005) identified five general mechanisms by which errors can be detected which include pathologist reviewing the case without additional information, with additional information, while preparing for a clinicopathologic conference, review on clinician’s request, or as a result of external consultation.Our present study was a post sign out audit which has both advantages and disadvantages.(Smith and Raab, 2012) Studies have noted higher error detection rates with retrospective reviews (Renshaw and Gould, 2006 Smith and Raab, 2012). Diagnostic disagreements in surgical pathology have ranged between 2.2% to 6.9% in some
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A look at Table 1 shows that misinterpretations of CNS lesions are the commonest errors made during Frozen Sections. Although we now see a large number of neopathology cases and some of us have subspeciality interests, difficult cases in every organ system are shown to the expert/s in that particular sub specialty as a mandatory second opinion before sign out. This regular practice on the whole works very well and acts as a safety valve.

However, on some occasions, few cases ‘slip through the net’ of this safety valve and may result in misinterpretations. Such lacunae or omissions in seeking mandatory second opinion in difficult cases are especially likely to occur in the summer month when a number of faculty members are on vacations. This lack of sub specialty practice will remain until there are sufficient surgical pathologists to allow each to practice within his/her specialized domain. A glance at tables 4 and 5 shows that many of the misinterpretations occurred as a result of failure to perform and examine deeper levels and/or special stains, failure to submit and examine adequate sections etc which point towards the problems discussed above. However, our results must be put in their proper perspective – misinterpretations occurred in 0.18% cases out of a total of 57,000 cases (including 0.36% of 2170 frozen sections).

Any changes made in our reports (after discovery of errors) are reported as “addendum reports”. However we use the term “amended report” when misinterpretations are involved. Even today, considerable confusion exists among pathologists regarding appropriate use of these designations especially if the changes are minor or “lesser changes”. However, most agree that a major change in diagnosis should be designated as an “amendment”. (Cooper, 2006; Nakhleh et al., 2006) A study by Finkelstein et al in 2012 reported that the number of cases with addendum reports increased from 0.9% in 1993 to as high as 8.6% in 2008 and that 5.6% of addendum reports had information which should have been reported as an “amended” rather than as “addendum” report. (Finkelstein et al., 2012)

The large majority of cases in our study were reviewed on the clinician’s request. Although a large Q-probe study of amended reports also showed that clinician requested reviews was the most common form of review, the percentage of such cases (20.5%) was very low compared to our study in which 90.5% cases were clinician requested reviews. (Nakhleh and Zarbo, 1998)

Omission of critical/important information was a major cause for errors in our surgical pathology reports accounting for 32.7% of all errors (Table 2). Many of the same reasons which are given above to explain the high percentage of misinterpretations can also be applicable to “errors of omissions”. However, we feel that the frequency of such errors in our practice has consistently declined especially since we switched to a synoptic /checklist format for the reporting of cancer resection specimens and will continue to decline in the future.

Typographic errors were also quite common in our setting (Table 2). Some of the causative factors include lack of English language proficiency of transcribers (and even residents), lack of synoptic /structured reporting format for many neoplastic lesions (specimens other than resections) and non-neoplastic lesions.

As described by Roy and Hunt, examination of amended reports (as in current study) is one of the various methods that are available for detecting errors in surgical pathology. (Roy and Hunt, 2010) The methods that we use for detection of errors include incedardepartmental consultation with one or more colleagues or intra-departmental consultation in the Departmental Consultation Conference which is held daily and in which multiple pathologists participate and give input on difficult and challenging cases. These methods detect errors before the case is signed out. Methods used for detecting errors after cases have been reported include diagnostic review for tumor boards, clincopathologic conferences etc. and when cases are reviewed at the time when a request for slides/blocks come from clinicians or patients for getting the case reviewed at another institution.

A look at Table 1 shows that misinterpretations of CNS lesions are the commonest errors made during Frozen Sections. Although we now see a large number of neopathology cases and some of us have subspeciality neuropathology interest and training (Ahmad et al., 2010) these errors occurred mainly in evening hours when a pathologist with subspeciality interest in neuropathology was not available.

A look at Table 7 shows that the greatest number of errors occurred in specimens from gastrointestinal tract (which also included liver, biliary tract and pancreas) followed by breast. We see a large number of cases from Gastrointestinal tract, liver and biliary tract and pancreas with extremely high rates of cancer in all these organs. (Ahmad et al., 2013; Ahmad et al., 2015) Specimens from lungs and other thoracic structures, lymph nodes and head and neck followed by specimens from urinary system and female genital tract were also quite prone to errors. Studies have also shown the propensity of gastrointestinal tract,
female genital tract, head and neck etc for errors.(Volmer et al., 2014; Chaudhary et al., 2014)

A percentage of interpretative errors in our series resulted at least partially from lack of relevant clinical information. It has been shown that 0.73% cases in surgical pathology require relevant clinical information for accurate diagnosis and a significant change in diagnosis occurs in over 60% such cases when the relevant clinical information is obtained,(Nakhleh et al., 1999) A good and alert clinician who reads a report carefully and in detail can be of great help in discovering errors in cases which have already been signed out.

Our findings were similar to other published studies. We need to develop documented procedures for timely review of cases to detect errors and monitor and document the result of these reviews and take steps to reduce errors.

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