Ophthalmic Service Guidance

Ophthalmic Pathology

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1. **What is ophthalmic pathology?**

Ophthalmic pathology is a laboratory-based discipline which provides an essential service to ophthalmologists and related professionals by examining samples taken from patients in order to provide a diagnostic opinion. Ophthalmic pathology is a subspecialty of cellular pathology (including histopathology and cytopathology), with much of its methodology based around light microscopic examination of glass slides on which are mounted sections of tissue or fluid preparations derived from the patient samples. These preparations are examined morphologically following use of staining methods including histochemistry and immunohistochemistry to highlight salient features of the specimen and aid in diagnosis. In some cases, ancillary methods such as molecular analysis or electron microscopy may be needed.

For the clinician, the "result" of an ophthalmic pathology specimen is generally a written report. Contents of the report of course vary according to the nature of the specimen and examination findings, but the report usually concludes with the reporting pathologist’s opinion on what the diagnosis is or, in less clear cut cases, suggestions as to possible diagnoses and what could be done (by the ophthalmologist or pathologist or some other party such as a geneticist) to achieve more clarity. For certain diagnoses, such as cancers, information may also be given about features such as tumour size or completeness of excision, in order to inform further management decisions.

2. **Where is ocular pathology provided?**

Reports on ophthalmic pathology specimens are provided either by consultant histopathologists and advanced practitioners who specialise fully in ophthalmic pathology, or by consultant histopathologists and neuropathologists who have a special interest in ophthalmic pathology as part of a broader diagnostic repertoire. Both groups of pathologists belong to the British Association for Ophthalmic Pathology (BAOP) and they all work within the structure of their own laboratories. There are some differences between the two groups in the provision of services described below.

In organisational terms, ophthalmic pathology services are provided through histopathology laboratories, with most dedicated ophthalmic pathology services being part of larger histopathology laboratories within hospitals. In England, ophthalmic pathology is nationally designated as a "highly specialised service" with central funding being provided to four ophthalmic pathology services (based in Liverpool, Manchester, Sheffield and London). These laboratories make up the National Specialist Ophthalmic Pathology Service (NSOPS), where the full time specialists in ophthalmic pathology are based. NHS specimens from England may be submitted to any of the NSOPS laboratories for examination (either primary reporting or second opinion) without charge to the referring organisation. There is a similar arrangement in Scotland. Details of the pathologists and laboratories which provide these services are available online.

Non-NSOPS BAOP members are pathologists who have eye pathology as a special interest, but who also have other areas of practice. NSOPS laboratories do not charge for NHS referrals, but non-NSOPS pathologists provide a more locally based service with tighter clinicopathological integration and responsiveness to local needs, in addition to having access to input from the centrally funded laboratories if desired.
3. Who should report ophthalmic pathology specimens?

Guidance provided by the Royal College of Pathologists states that pathologists reporting ophthalmic pathology specimens should participate in an appropriate external quality assurance (EQA) scheme. EQA participants usually receive a set of scheme-specific diagnostic cases twice a year for which they submit their responses (diagnoses) to the scheme organiser. Their responses are then scored against those of their peers, the other participants. Pathologists are expected to participate in schemes relevant to their area(s) of practice. For ophthalmic pathology, this is the Ophthalmic Pathology National EQA Scheme. NSOPS and BAOP pathologists and other Ophthalmic Pathology EQA Scheme participants are encouraged to attend the annual BAOP meeting, where the cases from the year’s scheme are discussed.

For certain types of tissue, it may be appropriate for the specimen to be reported by a pathologist with a different but related area of specialist pathology (who participates in the relevant EQA scheme), such as dermatopathology or neuropathology.

Reporting of specimens by pathologists with the appropriate specialist knowledge, skills and experience is good clinical practice and reduces risk of misdiagnosis as well as allowing the specialist pathologist to maintain and enhance his/her level of expertise. Additionally, it facilitates training for future specialists.

4. Why should the ophthalmologist send tissue to the laboratory?

It is not necessary for the ophthalmologist to send all tissue removed from every patient who has surgery, but there are published recommendations on what should be sent and what need not be sent. It can be useful for the ophthalmologist to correlate clinical diagnostic impression with histopathology findings. The major risk of discarding tissue is of missing a significant diagnosis (e.g. malignancy, epithelial downgrowth or sympathetic endophthalmitis), but it may be equally as valuable to know that removed tissue was histologically innocuous if the patient develops a subsequent lesion at the same site.

5. What specimens should the ophthalmologist send?

Much of the information below has been made available previously, in the joint guidance document of The Royal College of Ophthalmologists and The Royal College of Pathologists on referral of ophthalmic pathology specimens. The joint guidance document is currently available on an archived website and the information also forms part of an article in the RCOphth Focus Articles series. The information is presented here in tabulated form for ease of reference, but the content remains substantially the same.
<table>
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<tr>
<th>Specimens/surgery</th>
<th>What can be discarded</th>
<th>What to send</th>
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| Small lid biopsy       | • A first occurrence of a chalazion in a patient age >40 years  
                        | • A first or second occurrence of a chalazion in a patient age <40 years  
<pre><code>                    | • Excess skin removed at blepharoplasty, or normal tissue removed during a cosmetic procedure | All other material including recurrent chalazia (except as previously stated) |
</code></pre>
<p>| Full thickness eyelid  | Normal tissue removed during lid shortening, ectropion or entropion procedures          | All other material                                                        |
| Conjunctiva            |                                                                                        | All material should be sent including pterygia and pingueculae            |
| Cornea                 |                                                                                        | All material should be sent including Descemet's membranes                |
| Trabecular meshwork    | It is not necessary to send tissue removed at trabeculectomy                             | If there is a research interest, material may be sent by local agreement  |
| Iris, ciliary body, choroid | Peripheral iridectomy tissue from glaucoma or cataract surgery                          | All other material                                                        |
| Lens                   | It is not necessary to send material removed at cataract surgery                        | If there is a research interest, material may be sent by local agreement  |
| Vitreous               | It is not necessary to send intravitreal blood or opacities (e.g. asteroid hyalosis)    | Send material in cases where there is a suspicion of inflammatory disease (after bacterial samples have been taken) or malignancy (e.g. lymphoma) |
| Epiretinal membrane    | It is not necessary to send these unless there is clinical concern e.g. malignancy      | If there is a research interest, material may be sent by local agreement  |
| Subretinal membrane    | It is not necessary to send these unless there is clinical concern e.g. malignancy      | If there is a research interest, material may be sent by local agreement  |
| Evisceration and enucleation |                                                                                     | All of these should be sent for examination. There is a small but appreciable risk of there being an occult malignancy |
| Orbital biopsy         | It is not necessary to send normal soft tissue or bone removed during orbital decompression or squint surgery | All other tissue removed at surgery should be sent |</p>
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<th>What to send</th>
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<tbody>
<tr>
<td>Lacrimal gland</td>
<td></td>
<td>All removed tissue should be sent for examination</td>
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<tr>
<td>Lacrimal sac</td>
<td>It is not necessary to send bone removed during DCR</td>
<td>Lacrimal sac excisions should be sent for examination</td>
</tr>
<tr>
<td>Orbital exenteration</td>
<td></td>
<td>All of these should be sent for examination</td>
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<tr>
<td>Cytology</td>
<td></td>
<td>Impression cytology of the conjunctiva and corneal, and fine needle aspiration cytology of periocular or intraocular masses should be sent for examination. For aspirates of intraocular fluids see vitreous (above)</td>
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**Note:** This is not a prescriptive list. In any case where the surgeon has a clinical concern or question it is appropriate to send tissue for examination with a low threshold. Obviously, tissue taken for the purpose of histological/cytological diagnosis, such as a temporal artery biopsy, should be submitted for examination, even if it does not fall into one of the categories covered in the table.

Ophthalmologists are encouraged to discuss with their local BAOP pathologist or an NSOPS ophthalmic pathologist cases where there is uncertainty about whether/what to submit for examination.
6. Practical points

Communication
Communication with the laboratory is key. If there is clinical urgency, if the clinician is unsure how to handle a particular type of specimen, or if there may be a need for non-routine handling of the specimen (such as fresh material or electron microscopy), the laboratory should be contacted in advance.

Request forms
Histopathology/cytology request forms vary between hospitals, but include standard information fields. It is therefore acceptable for the ophthalmologist to use a request form available locally, even if the specimen is to be submitted to a laboratory elsewhere for examination. A downloadable request form for the NSOPS laboratories is available. The request form should be completed fully including the following:

- Patient details: Surname, forename, date of birth, hospital number, NHS number
- Clinician details and hospital location. A telephone number is useful if clarification is needed, or if there is a need for urgency
- Date specimen taken
- High risk status
- Specimen type
- Relevant clinical information

Fixation and containers
Patient identification and specimen details should be completed on each specimen pot submitted. Multiple specimens from the same patient should be placed in different individually labelled containers to avoid confusion. Unless agreed otherwise in advance (see "Unfixed specimens" below), specimens should be fixed in 10% neutral buffered formalin. **This includes suspected sebaceous carcinoma specimens which do not need to be submitted fresh.** The volume of fixative (and therefore size of specimen pot) should be appropriate to the size of the specimen. As a rule of thumb, 10x the specimen volume of formalin is sufficient for adequate fixation.

Specimen packaging and transport
Specimens which are not dealt with by the ophthalmologist’s hospital laboratory service will usually be sent to the examining laboratory by post or courier. Packaging of diagnostic specimens (usually classified as Category B biological substances) must conform with the UN3373 and P650 packaging instruction. Further information is available from the postal service or courier selected. It is recommended that fax/phone back arrangements are implemented to ensure confirmation of receipt by the laboratory. It may be easier for specimens to be sent via the ophthalmologist’s local laboratory rather than directly from the ophthalmology department.

Unfixed specimens
Rarely, it is appropriate to submit unfixed material for examination, e.g. for frozen section examination or immunofluorescence. These specimens should only be submitted following prior arrangement with the laboratory including contact details of the responsible clinician, so that any late or non-arrival of the specimen may be notified to the clinician. Specimens must be delivered without delay in order to allow handling in a timely fashion. Conjunctival specimens for immunofluorescence may be stabilised for transport in Michel’s medium.
Note: High Risk specimens should never be submitted unfixed

Rapid processing specimens
Many histopathology laboratories provide a service for intraoperative diagnosis (frozen section) and/or rapid paraffin processing (for delayed reconstruction). These services are provided by local arrangement, and ophthalmologists should discuss their requirements with their chosen laboratory.

Turnaround times
The Royal College of Pathologists Key Performances Indicator recommendations states that 90% of diagnostic biopsies should be reported within 7 days of biopsy. This figure does not take into account transit times between the ophthalmology department and the reporting laboratory. Individual laboratories can provide information on their current turnaround times on request.

Research
Sometimes the ophthalmologist wishes to sample ocular tissue for research. If the tissue is also potentially of diagnostic significance, it is recommended that the ophthalmologist communicates with the pathologist who will be handling the specimen for diagnostic purposes, even if the pathologist is not involved in the research project.

Awareness of errors
As previously stated, the histopathology report provides the pathologist's opinion on the submitted specimen. There is potential for mistakes to occur at any point of the specimen's pathway, including misidentification of the patient or specimen, technical malfunctions, and errors in diagnosis by the pathologist. If the clinical impression and pathology report do not "fit" with each other, it is better for the ophthalmologist to discuss this with the pathologist sooner rather than later.
7. Reading

5. Focus Articles. Available at: https://www.rcophth.ac.uk/standards-publications-research/focus-articles/ (Accessed: 8th March 2016)

8. Author

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