EVEREST study report 2: imaging and grading protocol, and baseline characteristics of a randomised controlled trial of polypoidal choroidal vasculopathy

Colin S Tan,1,2 Wei Kiong Ngo,2 Jian Ping Chen,2 Nikolle W Tan,1,2 Tock Han Lim,1,2 on behalf of the EVEREST Study Group

ABSTRACT

Purpose To describe the imaging standards, grading protocol and baseline characteristics of polypoidal choroidal vasculopathy (PCV) from the EVEREST study.

Methods In a prospective, multicentre study, confocal scanning laser ophthalmoscope indocyanine green angiography (ICGA) was performed using a standardised imaging protocol. All images were graded using standardised, calibrated equipment by fellowship-trained ophthalmologists at the Central Reading Center.

Results Sixty-one patients with PCV were included in the study. ICGA characteristics included: nodular appearance stereoscopically (56 eyes, 91.8%), hypofluorescent halo (42, 68.9%), abnormal vascular network (54, 88.5%) and pulsation of the polyps (4, 6.6%). Colour fundus photography revealed orange subretinal nodules (34, 55.7%) and massive submacular haemorrhage (8, 13.1%).

Conclusions The imaging and grading protocols and baseline characteristics of a multicentre, randomised controlled trial of PCV are described in detail, and may serve as reference for future randomised, controlled trials on PCV.

METHODS
This study was approved by the respective institutional review boards of the study sites. Written, informed consent was obtained from all patients prior to enrolment in the study.

Imaging protocol
Fluorescein angiograms and ICGA were performed using a confocal scanning laser ophthalmoscope (Heidelberg Retinal Angiograph (HRA), HRA-C/HRA2/HRA Spectralis, Heidelberg Engineering, Germany). When performing confocal scanning laser ophthalmoscope angiograms, the dose of indocyanine green (ICG) required is significantly lower compared with flash ICGA. ICG 10 mg in 2 mL using the accompanying aqueous diluent kit (Akorn, Pulsion or equivalent, as approved by the local health authority) was injected through the anterior cubital fossa vein over ≤5 s.

The standard angiographic field for all frames was 30° centred on the fovea, with a resolution of...
1536×1536 pixels and capturing speed of ≥6 frames per second for dynamic angiography. For still-frames, image quality was improved by digital signal averaging using either the automated real-time function or the 'mean' function.

The timer was started at the start of the bolus injection, while the capture button was triggered only at first dye appearance. The 'precapture' mode of the HRA device allows recording of 2 s of the angiogram prior to the triggering of the capturing button, which is very useful for capturing the initial frames when the dye just enters the choroidal circulation through the feeder vessel. In order to image the full filling sequence of the branching vascular network (BVN), the dynamic capture duration was set at ≥30 s.

Still-frame captures were performed according to the sequence as shown in online supplementary appendix 1. Stereo-pair images were obtained during each time period. To obtain stereo-pair images, one frame was taken (using the automated real-time function) and the device was shifted laterally within the same plane, slightly to the right, and the second stereo-pair image taken.

Case definition of PCV

The Central Reading Center used standardised criteria to diagnose PCV, the features of which were distilled from available English literature from 1990 to 2008. PCV was diagnosed based on early subretinal ICGA hyperfluorescence (appearing within the first 5 min of ICG dye injection) and at least one of the following diagnostic criteria (figure 1):

- Nodular appearance of the polyp on stereoscopic viewing
- Hypofluorescent halo around the nodule
- Abnormal vascular channel(s) supplying the polyps
- Pulsatile filling of polyps
- Orange subretinal nodules corresponding to the hyperfluorescent area on ICGA
- Massive submacular haemorrhage.

Two types of abnormal vascular channels were identified using dynamic ICGA (figure 2). A typical BVN originates from a feeder vessel and spreads outwards, with polyps typically located at the periphery. A second, less common type was interconnecting channels, which consisted of a network of criss-crossing choroidal vessels which supply the associated polyps. The dye has no specific point of origin or direction of flow, unlike a BVN.

Grading protocol

All images were graded by trained graders (Fellowship-trained retinal specialists) at the Fundus Image Reading Center, National Healthcare Group Eye Institute, Singapore, using standardised equipment and grading protocols. High resolution (3000×2000) monitors (HP LP3065, Hewlett Packard, Palo Alto, California, USA) were used to view all images for stereoscopic viewing at the full resolution of the ICGA image. The monitors were calibrated weekly using the Spyder 3 calibration software (Datacolor, Lawrenceville, New Jersey, USA).

ICGA images were graded using the Heidelberg Eye Explorer software (Heidelberg Engineering, Heidelberg, Germany). The dynamic ICGA was played for the entire 30 s. The slider on the viewing software was then moved to the point when the ICG dye first appeared and advanced slowly to allow visualisation of the initial filling of the feeder vessels and polyps. In addition, the pulsation of the polyps was determined using the dynamic ICGA.

The filling times were read from the timer of the viewing software. Vascular channel filling time was defined as the first time the ICG dye was visible within the abnormal vascular channels. Polyp filling time was defined as the time of first dye appearance within a polyp.

A best-fit circle was drawn around each polyp using the drawing tools on the Heidelberg Eye Explorer software (figure 3A). The diameter of each circle was used to calculate the area of each polyp. The total lesion area was determined using the freehand drawing tool. The grader drew an outline which encompassed all polyps identified and the area of the abnormal vascular channels (identified using dynamic ICGA), and the software returned the total area in mm². The circle drawing tool was then used to draw a best-fit circle around the total lesion area, the diameter of which was the greatest linear dimension (figure 3B).

Stereo-pair colour fundus photographs were viewed using the Visupac viewer V4.4.4 (Carl Zeiss Meditec, Dublin, California, USA) and the Screen-Vu stereoscope. The presence of orange subretinal nodules and massive submacular haemorrhage (defined as submacular haemorrhage ≥4 disc areas) was noted from the colour fundus photographs.

RESULTS

Of 95 cases submitted for screening, 61 were diagnosed with PCV. The mean age was 65 years (range, 46–84 years, SD ±9.2 years), with 41 men (67.2%) and 20 women (32.8%). All patients were of Asian ethnicity. In 32 patients, the right eye was involved (53.5%) whereas the left eye was involved in 29 patients (47.5%).

The frequency of ICGA characteristics were as follows: nodular appearance, 56 (91.8%); hypofluorescent halo around the nodules, 42 (68.9%); presence of abnormal vascular network, 54 (88.5%); and pulsation of the polyp(s) on dynamic ICGA, 4 (6.6%). Using colour fundus photography, 34 (55.7%) had orange subretinal nodules and 8 (13.1%) presented with massive submacular haemorrhage.

The characteristics of the polyps and BVN are summarised in table 1. The number of polyps ranged from 1 to 9, with a median of 4. The polyp configurations included solitary—16 (26.2%), ring—28 (45.9%), cluster—15 (24.6%) and mixed configurations—2 (3.3%). The mean total polyp area was 0.25 mm² (median 0.21 mm²; range, 0.025–2.11 mm², SD ±0.28 mm²). The area of the PCV lesion (comprising polyps and associated vascular channels) ranged from 0.2 mm² to 10.7 mm², with a mean of 3.11 mm² and a median of 2.28 mm².

Of the cases with associated vascular channels, 46 (75.4%) had a distinct BVN, whereas 8 (13.1%) were classified as interconnecting channels, as described above. The vascular channels (either BVN or interconnecting channels) filled within 7.3–32.0 s, with a mean of 17.9 s (median: 17.6 s) while the mean filling time for the polyps was 21.9 s (median: 22.0 s; range, 7.3–40.4 s, SD±8.1 s).

The patient manifested with a variety of clinical presentations, including serous subretinal fluid (28 (45.9%)), serosanguinous (23 (37.7%)), haemorrhagic (2 (3.3%)) and massive submacular haemorrhage (8 (13.1%)). Patients with massive submacular haemorrhage were less likely to have abnormal vascular channels on ICGA (37.5% vs 83.0% for those without massive haemorrhage, p=0.001). A higher percentage of patients with pulsation of the polyp on dynamic ICGA presented with haemorrhage of any size, 1 of 4 (25.0%) compared with 9 of 57 (15.8%) for those without pulsation, but the difference was not statistically significant (p=0.63). The frequency of
haemorrhagic presentations did not differ between those with orange subretinal nodules (5 patients, 14.7%) and those without nodules (5 patients, 18.5%).

DISCUSSION
In this paper, we have described the ICGA imaging standards, grading methods and diagnostic criteria used in the EVEREST Study, the first multicentre randomised controlled trial of PCV treatment. The baseline ICGA and colour fundus photograph characteristics of patients with PCV are also described. These methods may provide the reference for standardised imaging and grading protocols for PCV in future multicentre studies.

The relevance of the diagnostic criteria can be seen from the results of this paper. While earlier studies on PCV used a variety of clinical and ICGA features to diagnose PCV and differentiate it from neovascular AMD, these features were highly variable between studies and the characteristic appearance of a polyp was often not well-defined. In many papers, the polyps have been described as ‘polypoidal choroidal vascular dilatations’, ‘focal vascular dilations’, or ‘polypoidal lesions’. These terms, however, may mean different things and be interpreted differently by various clinicians. Therefore, it is imperative to use standardised criteria that are easy to apply across different studies in order to ensure that the results of these studies are comparable. In this study, three parameters were
used to describe a typical polyp using ICGA—nodular hyperfluorescence on stereoscopic ICGA (91.8%), hypofluorescent halo around the nodule (68.9%) and pulsation during dynamic ICGA (6.6%). We believe that stereo-pair ICGA images are essential in the diagnosis of PCV, as it allows the determination of nodularity (the most common diagnostic feature) and the depth of the lesions, which helps to differentiate vascular polyps from retinal lesions such as macroaneurysms, large microaneurysms and retinal angiomatous proliferation. This was of particular importance because spectral domain optical coherence tomography (SD-OCT) was not widely available in 2008 when this trial was underway. In future studies, SD-OCT may play an important role in determining the depth and shape of the PCV lesion.

Dynamic ICGA is an essential component of PCV diagnosis, as it allows identification of two of the six diagnostic criteria. The exact boundaries of abnormal vascular channels of the PCV lesion are best visualised in the early phase of the angiogram (within 30 s) and may not be seen as clearly after 1 min due to complete filling of the normal choroidal vessels. In addition, the presence of pulsation cannot be detected without the use of dynamic ICGA.

The abnormal vascular channels began to fill within 7.3–32 s from injection. Therefore, the use of dynamic ICGA for the first 30 s is sufficient to detect these vessels. In all cases, the polyps started to fill within 40.4 s from injection, with a mean of 21.9 s, which corresponds to the original description of early, intense focal hyperfluorescence on ICGA. Therefore, the use of either the 1-min or 3-min stereo-pair ICGA images should be adequate to assess the presence, location and extent of polyps and a later stereo-pair at ≥10 min may not be necessary. True late-onset focal hyperfluorescences (appearing after 6 min) are more likely to be focal ICG staining of unhealthy RPE or choroidal vascular knuckle showing through focal RPE atrophy (window defect). In both cases, stereo viewing would help to differentiate them.

In this series, 33 of 61 cases (54.1%) presented with subretinal haemorrhage. This highlights the importance of haemorrhage as diagnostic criteria for PCV as has previously been described. Ahuja et al reported that 85% of patients with haemorrhagic and exudative detachments had evidence of PCV on ICGA.

In several earlier papers, the presence of orange subretinal nodules was used to diagnose PCV. This clinical feature was seen in 55.7% of cases in this study, thus highlighting that not all cases of PCV manifest with characteristic nodules and the need for ICGA to identify the PCV lesion.

The strengths of this study include the use of standardised imaging protocols and equipment on all cases of PCV. The diagnosis of PCV was made by applying stringent diagnostic criteria by expert graders from the Central Reading Center. The main limitation is the lack of data from OCT, which can potentially be used as one of the diagnostic criteria. In future studies, SD-OCT can play an invaluable role in identifying features which are characteristic of PCV and confirming the location of the hyperfluorescent nodule.

In conclusion, we have described the standardised imaging protocols, grading methods and diagnostic criteria used to diagnose PCV and highlighted important features in a prospective series of patients with PCV. It is hoped that these detailed techniques, and the baseline characteristics described, can be useful to clinical practice and future randomised controlled trials on PCV.

**Collaborators** Members of the EVEREST Study Group: (1) Adrian Koh, MD FRCS, National Healthcare Group Eye Institute, Tan Tock Seng Hospital, Singapore; (2) Won Ki Lee, MD PhD, Seoul St Mary’s Hospital, The Catholic University of Korea, Seoul, Korea; (3) Lee-Jen Chen, MD, Mackay Memorial Hospital, Taipei, Taiwan; (4) Shih-Jen Chen, MD PhD, Taipei Veterans General Hospital, Taipei, Taiwan; School of...
REFERENCES
Appendix 1. Fluorescein and indocyanine green angiography imaging sequence.

**Dynamic capture** (video angiography) for 30 seconds after the first appearance of the dye in the retinal (fluorescein angiography) or choroidal circulation (indocyanine green angiography).

**Still-frame 30-degree stereo pair images using automated real time setting** as follows:

1 min – 1 min 59 sec: 1 stereo pair of *study* eye

2 min – 2 min 59 sec: 1 stereo pair of *fellow* eye

3 min – 3 min 59 sec: 1 stereo pair of *study* eye

4 min – 4 min 59 sec: 1 stereo pair of *fellow* eye

5 min – 5 min 59 sec: 1 stereo pair of *study* eye

10 min - 10 min 59 sec: 1 stereo pair of *study* eye

11 min - 11 min 59 sec: 1 stereo pair of *fellow* eye

20 min - 20 min 59 sec: 1 stereo pair of *study* eye

21 min - 21 min 59 sec: 1 stereo pair of *fellow* eye