

# Fluorescence Imaging • Fluorescein angiography • ICG angiography • Fundus autofluorescence

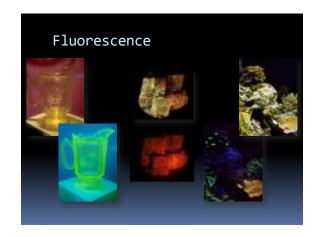
## Fundus Autofluorescence (FAF)

- Fundus autofluorescence (FAF) is a diagnostic imaging technique for documenting the presence of fluorophores in the human eye.
- Fluorophores are chemical structures that possess fluorescent properties when exposed to electromagnetic energy of an appropriate wavelength.

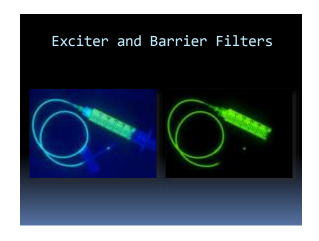
## Fluorescence

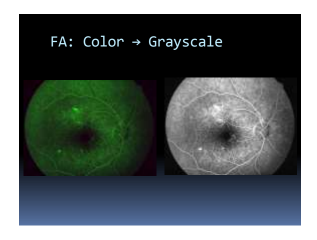
- When fluorophores absorb light of a particular wavelength, they are temporarily excited to a higher energy state.
- Triggers the emission of light at wavelengths longer than the excitation source.

# Fluorescence • Emission occurs only as long as the fluorescent subject remains illuminated by the exciting source (10<sup>-8</sup> seconds).



# Fluorescein Absorbs blue light, with peak absorption and excitation occurring at wavelengths between 465-49onm. Fluorescence occurs at the yellow-green wavelengths of 520 to 530nm.







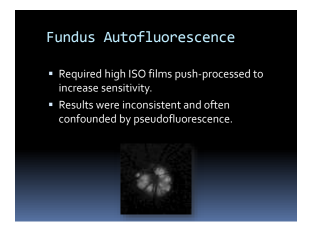
## Fundus Autofluorescence

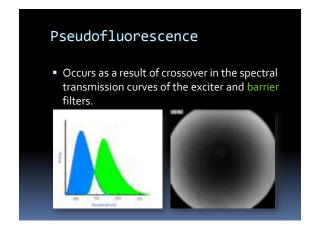
- The term "autofluorescence" is used to distinguish fluorescence that can occur naturally vs. fluorescence that is derived from administration of fluorescent dyes.
- Optic nerve drusen, astrocytic hamartomas, lipofuscin pigments in the retina, and the aging crystalline lens are all believed to exhibit natural fluorescence.

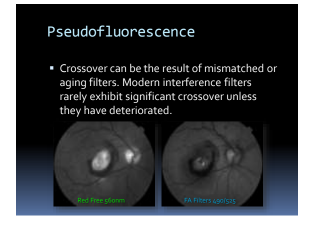
## Fundus Autofluorescence

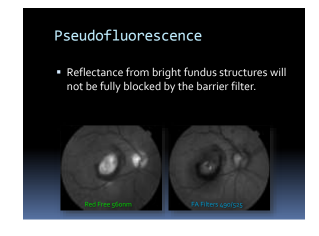
 Procedures for documentation of highly fluorescent entities such as optic disc drusen have been employed for years with varying degrees of success using a fundus camera with fluorescein excitation and barrier filters.

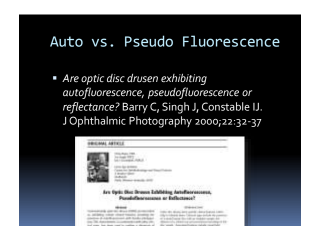


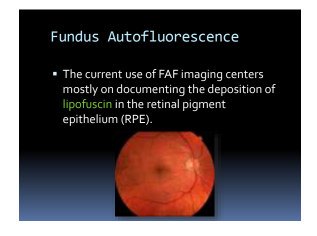












## Renewed Interest in AF

 Autofluorescence imaging of lipofuscin first became practical with the implementation of confocal scanning laser technology.



### Fundus Autofluorescence

- Lipofuscin is a fluorescent pigment that accumulates in the RPE as a metabolic byproduct of cell function.
- Lipofuscin deposition normally increases with age, but may also occur from RPE cell dysfunction or an abnormal metabolic load on the RPE.

## Fundus Autofluorescence

- There are as many as ten different fluorophores found in lipofuscin.
- The dominant fluorophore in lipofuscin is A2-E, a compound consisting of two vitamin A molecules and ethanolamine.
- A2-E possesses toxic properties that may interfere with normal RPE cell function.

### Fundus Autofluorescence

- FAF imaging is particularly challenging due to low levels of fluorescence and variability in the amount of lipofuscin present depending on age, health of the RPE, and disease process.
- Requires a more light-efficient method than the traditional technique used for imaging disc drusen.

## Fundus Autofluorescence

- There are two different digital technologies currently used to capture fundus autofluorescence images:
  - cSLO
  - Modified fundus camera.





## Fundus Autofluorescence

- Both systems require significant amounts of light and high gain settings to achieve adequate exposure, and are susceptible to unwanted noise that can interfere with image detail.
- Noise is false pixel data that occurs from poor signal-to-noise ratios and the amplification needed to record fluorescence.

## Scanning Laser Ophthalmoscope

 The confocal scanning laser ophthalmoscope (cSLO) is an instrument that can be used for several retinal imaging modalities including fluorescein angiography, ICG angiography and fundus autofluorescence.



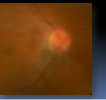


## Scanning Laser Ophthalmoscope

- A monochrome laser scans across the fundus in a raster pattern to illuminate and record successive elements of the retina, point-bypoint at speeds up to 24 milliseconds.
- The laser delivers a very narrow wavelength band allowing for efficient excitation of fluorescence.

## Scanning Laser Ophthalmoscope

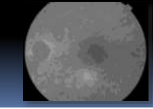
 A confocal aperture positioned conjugate to the focal plane of the retina blocks non image-forming light from reaching the sensor to minimize scatter and improve contrast.

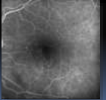




## Scanning Laser Ophthalmoscope

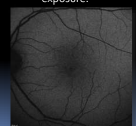
 Confocal imaging reduces the effects of short wavelength scatter in the ocular media and confounding AF from the crystalline lens.



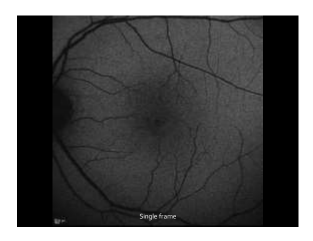


## cSLO FAF Sampling

Sampling smoothes noise and increases exposure.

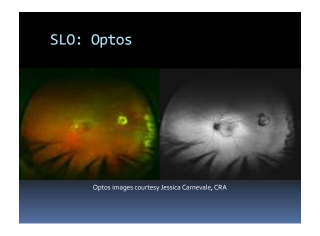


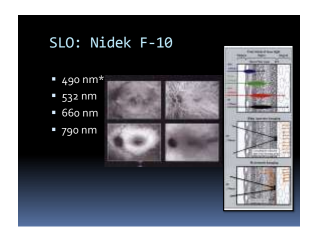


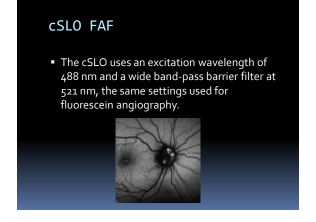




## SLO: Spectralis HRA FA excitation and blue reflectance (red free) 488nm solid state laser ICG excitation 790nm diode laser IR Reflectance 820nm diode laser



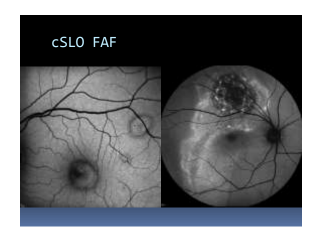




## FAF imaging must be done before angiography if both procedures are performed with a cSLO on the same visit. Even the slightest amount of intravenous.

cSLO FAF

- Even the slightest amount of intravenous fluorescein will compromise the effectiveness of cSLO FAF.
- Residual topical fluorescein may also interfere with FAF.



## Modified Fundus Camera FAF

 More recently (2003), digital fundus-camera based systems have been developed for autofluorescence imaging.

Fundus Autofluorescence and Age-related Macular Degeneration

Billed F. Sosk, 347

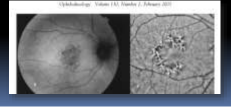
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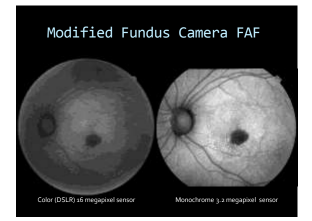
## Modified Fundus Camera FAF

 Utilizes high-sensitivity monochrome digital sensors with different filter combinations than used for angiography.



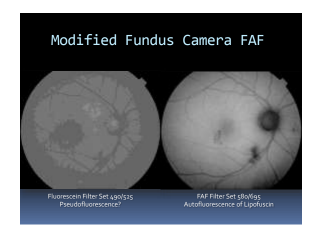
## Modified Fundus Camera FAF

- Monochrome digital backs are generally considered better than color sensors for retinal angiography and autofluorescence.
- They are more light-sensitive, and all pixels are available for exposure by the relatively limited band of wavelengths generated by fluorescence.

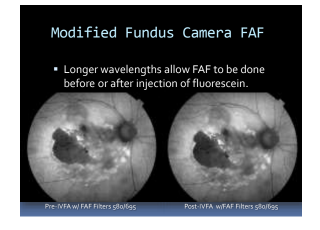


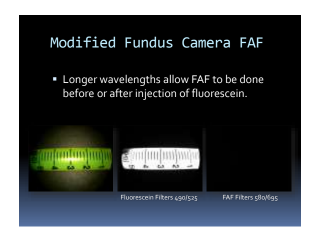
## Modified Fundus Camera FAF

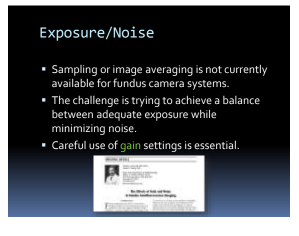
- The digital fundus camera technique first described by Spaide employs an excitation filter centered at 580 nm and a barrier filter centered at 695 nm.
- These wavelengths are shifted toward the red end of the spectrum to avoid unwanted short-wavelength autofluorescence from the crystalline lens.

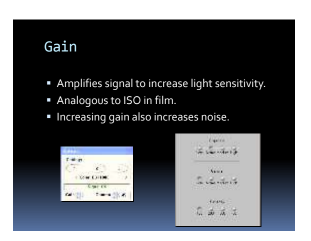


# FAF imaging can be conducted either before or after fluorescein angiography with fundus camera based systems. The FAF excitation wavelength of 580 nm causes minimal excitation of fluorescein and the barrier filter centered at 680 nm blocks the emission peak of fluorescein (520 nm).



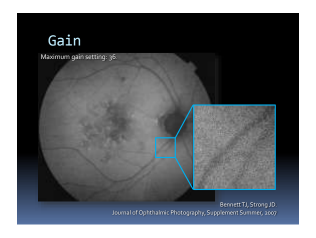


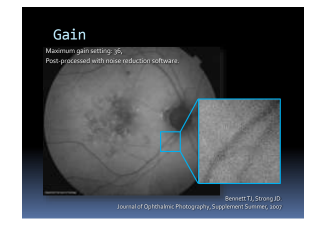


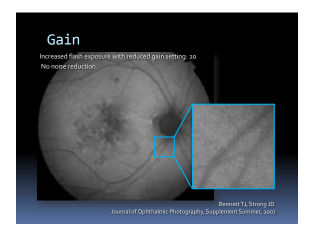








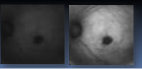


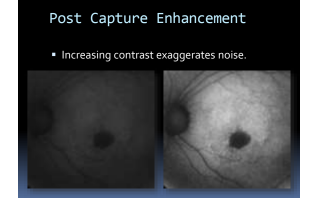


# Modified Fundus Camera FAF The default camera controls for FAF typically place the gain near the maximum setting in order to record low-level fluorescence. There may be very little room for lowering gain to reduce amplifier noise and still maintain sufficient exposure.

## Modified Fundus Camera FAF

- If the gain setting is too low, underexposure can occur resulting in dark, low-contrast photographs.
- Enhancement of underexposed images to improve brightness and contrast will increase noise in a manner similar to increasing gain.





## Gain

- In very low-light situations where high gain is used, like fundus autofluorescence:
  - Set the fundus camera for maximum light transmission and flash output.
  - Control exposure using gain to ensure lowest possible gain setting to reduce noise.

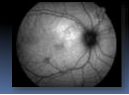


## Maximizing Transmission

- All controls should be set for maximum light transmission and flash output.
- Light transmission may be best at the widestangle setting in some variable angle fundus cameras.
- If the fundus camera is equipped with an illumination diaphragm it should be set to the largest aperture.

## Maximizing Transmission

 When light transmission is maximized, eyes with significant accumulation of lipofuscin can be imaged with reduced gain settings while still maintaining adequate exposure.



## Maximizing Transmission

- In the absence of significant accumulation of lipofuscin, underexposure can still occur in some widely dilated eyes with clear media.
- Young patients.



## New "Spaide" Filters

- New, more efficient filter sets that significantly improve light transmission are now commercially available for some funduscamera based systems.
- The patented Spaide filters are only available for Topcon Imagenet systems, but other manufacturers may also have 2<sup>nd</sup> generation filter sets.

## New "Spaide" Filters

- The excitation filter has a band-pass range of about 535-585 nm and the barrier filter has a band-pass range of about 605-715 nm.
- Avoids excitation of both the crystalline lens and fluorescein, improves light transmission, and reduces noise.
- FAF imaging can be done either before or after fluorescein angiography.

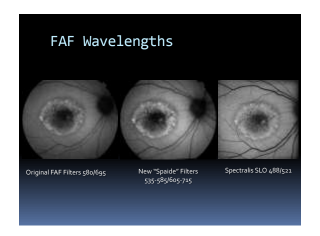
## New "Spaide" Filters

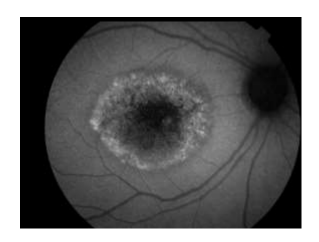
 The more recent generation of filter sets dramatically improve light transmission, allowing for lower gain settings and improved exposure.

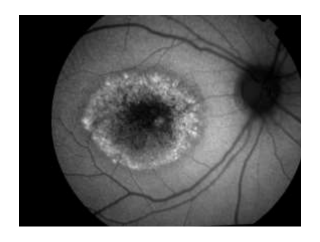


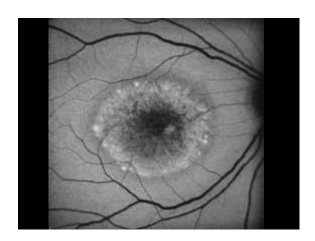
# New "Spaide" Filters Original FAF Filters: Gain 36, 300ws New Spaide Filters: Gain 16, 300ws

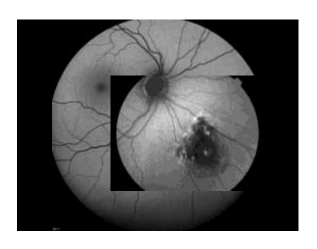
# CSLO Exciter: 488 nm Barrier: 521 nm (short cutoff/wide bandpass) Original Spaide filters: Exciter: 580 nm Barrier: 695 nm New proprietary "Spaide" filters: Exciter: 535-585 nm Barrier: 605-715 nm

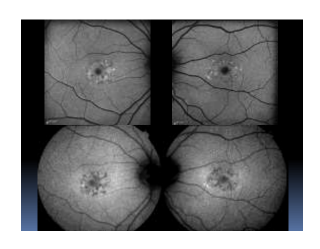












## Instrument/Camera Technique

- Proper camera technique is necessary to obtain quality FAF images.
- Target the focus to the level of the retinal pigment epithelium, which is where the majority of autofluorescence typically occurs.
- Once images are captured at this focus level, the camera can be refocused on different layers when autofluorescence is detected in other retinal structures.

## Instrument/Camera Technique

- cSLO: If viewing in IR prior to FAF, focus will need to be adjusted to account for the shorter wavelengths of the blue laser (488 vs 820 nm).
- Turn the focus knob approximately ¼ turn clockwise.





## Instrument/Camera Technique

- Exposure can be improved with optimal axial alignment of the illuminating beam within the center of the dilated pupil.
- The ability to move the beam within the pupil to avoid prominent media opacities also helps.
- Maximum pupillary dilation will allow even illumination and exposure.

## FAF Findings

- The optic nerve, retinal blood vessels, and the fovea normally appear dark against a variable background of fluorescence from the RPE.
- The absence of the RPE at the optic nerve head causes it to appear dark.



## FAF Findings

 Retinal vessels block both the excitation and emission of fluorescence from the underlying RPE and also appear dark.



## FAF Findings

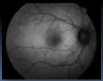
 The density and morphology of pigment in the fovea causes absorption of the excitation wavelengths making the fovea appear darker than the surrounding macula, especially with the cSLO.

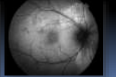




## Findings/Interpretation

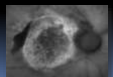
 Autofluorescence imaging is effective because it can detect metabolic changes in a cell monolayer, the retinal pigment epithelium, making it useful in conditions where the health of the RPE plays a key role.





## Findings/Interpretation

- Hyperfluorescence is a sign of increased lipofuscin accumulation, which may indicate degenerative changes or oxidative injury.
- Hypofluorescence usually indicates missing or dead RPE cells.

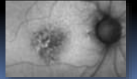


## Documentary vs. Diagnosis

- Documentary:
  - Geographic atrophy
  - Pigmentary changes (RP, ICSC...)
- Diagnostic:
  - Early detection of bullseye/retinal toxicity
  - Progression of geographic atrophy
  - Buried disc drusen
  - Macular hole
  - ICSC activity/leakage?

## Diagnostic Applications

 The role of lipofuscin in the pathogenesis of macular degeneration is currently unknown, but increased autofluorescence may precede development or progression of geographic atrophy in ARMD.



## Diagnostic Applications

 Geographic atrophy that appears as a "window defect" in fluorescein angiography will appear dark in autofluorescent imaging.

