Detecting Tree-like Multicellular Life on Extrasolar Planets

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Abstract

Over the next two decades, NASA and ESA are planning a series of space-based observatories to find Earth-like planets and determine whether life exists on these planets. Previous studies have assessed the likelihood of detecting life through signs of biogenic gases in the atmosphere or a red edge. Biogenic gases and the red edge could be signs of either single-celled or multicellular life. In this study, we propose a technique with which to determine whether tree-like multicellular life exists on extrasolar planets. For multicellular photosynthetic organisms on Earth, competition for light and the need to transport water and nutrients has led to a tree-like body plan characterized by hierarchical branching networks. This design results in a distinct bidirectional reflectance distribution function (BRDF) that causes differing reflectance at different sun/view geometries. BRDF arises from the changing visibility of the shadows cast by objects, and the presence of tree-like structures is clearly distinguishable from flat ground with the same reflectance spectrum. We examined whether the BRDF could detect the existence of tree-like structures on an extrasolar planet by using changes in planetary albedo as a planet orbits its star. We used a semi-empirical BRDF model to simulate vegetation reflectance at different planetary phase angles and both simulated and real cloud cover to calculate disk and rotation-averaged planetary albedo for a vegetated and non-vegetated planet with abundant liquid water. We found that even if the entire planetary albedo were rendered to a single pixel, the rate of increase of albedo as a planet approaches full illumination would be comparatively greater on a vegetated planet than on a non-vegetated planet. Depending on how accurately planetary cloud cover can be resolved and the capabilities of the coronagraph to resolve exoplanets, this technique could theoretically detect tree-like multicellular life on exoplanets in 50 stellar systems.

Key Words: TPF—Darwin—Extrasolar planets—Astrobiology—BRDF—Remote sensing—multicellular life.

1. Introduction

Over the next few decades, NASA and ESA are planning a series of space-based observatories to measure the spectra from Earth-sized extrasolar planets in regions where liquid water may be present (Lawson et al., 2007; Cockell et al., 2009). There has been much justifiable interest in developing methods of detecting life on these planets. Planetary biosignatures that have been considered include biogenic gases in the atmosphere (O₂ in the presence of H₂O, O₂, CH₄ with O₂, CH₃Cl, N₂O) (Des Marais et al., 2002) and surface reflectance spectra of vegetation (Ford et al., 2001; Seager et al., 2005; Tinetti et al., 2006a, 2006b; Kiang et al., 2007a, 2007b).

A potential biosignature is a planet’s absorption spectrum (Kiang et al., 2007b). Earth’s plants have a sharp, order-of-magnitude increase in leaf reflectance between 700 and 750 nm wavelengths due to the difference between photosynthetic absorption of photons (between 400 and 700 nm) and reflectance and scattering at higher wavelengths not used for photosynthesis (>700 nm). Seven hundred nanometers is in the red part of the electromagnetic spectrum, and the reflectance contrast is referred to as the “red edge”. Such a sharp, order-of-magnitude change in reflectance is rare in nature, and Earth’s red edge has been confirmed as a planetary biosignature both in the spectrum of Earthshine (spatially integrated light scattered off the Moon and reflected to Earth) (Seager et al., 2005; Palle et al., 2009) and by remote sensing from the Galileo spacecraft (Sagan et al., 1993). Researchers have theorized that water-splitting photosynthesis is the only type of photosynthesis efficient enough to support plentiful life on an extrasolar planet (Wolstencroft and Raven, 2002). The wavelength of an edge-like reflectance feature like the red edge could theoretically be determined if the planet’s distance from the Sun and the photons required to split water at this distance are known (Wolstencroft and Raven, 2002; Kiang et al., 2007b).

If future space observatories such as Terrestrial Planet Finder (TPF; NASA) or Darwin (ESA) succeed in finding a
planet with both biogenic gasses in the atmosphere and a red-edge characteristic of life, we would want to know whether the land-based life on this planet is similar to the green photosynthetic slime common for most of Earth’s history (Beerling, 2007) or whether this life has evolved to be diverse and multicellular. For instance, on Earth, the first fossil evidence of land plants, similar to modern bryophytes, was in the Mid-Ordovician (490–430 million years ago) (Graham et al., 2000). Single-celled photosynthetic organisms evolved as early as 3 billion years ago and were present on land as early as 1.2 billion years ago (Horodyski and Knauth, 1994). High $\delta^{13}C$ values in Precambrian rocks indicate that there was a significant cover of single-celled photosynthetic organisms on land during this period (Kenny and Knauth, 2001). Therefore, viewed from space, from 1.2 billion years ago to 490 million years ago, the Earth could have had a red-edge signal and biogenic gases in the atmosphere but no visible multicellular life. Below, we explore the question of whether multicellular life is detectable and whether vegetated planets can be distinguished from planets with less-complex life.

Multicellular life on Earth is characterized by hierarchical branching networks, a topological structure ubiquitous in biology (Brown and West, 2000). For multicellular photosynthetic organisms, this branching network is characterized by a “tree,” whose structural attributes and fractal topology are conserved over different phyletic groups and are independent of the environmental conditions that accompany ontogeny (West et al., 1999). Competition for light and the need to transport water and nutrients in multicellular photosynthetic organisms has led to the tree-like structure on Earth that emerges from a few general principles that are widespread in nature (West et al., 1999). As a consequence of these biomechanical and evolutionary constraints, the tree growth habit has evolved independently several times on Earth (Donoghue, 2005). We make the assumption that such biomechanical constraints combined with Darwinian evolution will likewise produce tree-like structures for photosynthetic multicellular organisms, if they exist, on extrasolar planets.

The same constraints that result in tree-like shapes of individual organisms result in predictable relationships between size, shape, and population density at the stand-level in forests (Enquist et al., 1998; West et al., 2009). These variations in the size and shapes of trees in forest stands result in strong differences in the reflectance of forest canopies as seen from different angles overhead (Li and Strahler, 1992; Bréon et al., 1997, 2002; Wolf et al., 2010).

The bidirectional reflectance distribution function (BRDF) is the change in observed reflectance with changing view angle or illumination direction (Schaepman-Strub et al., 2006). The BRDF is the ratio of the differential radiance (dL) (W m$^{-2}$ sr$^{-1}$) to the differential irradiance (dE) (W m$^{-2}$), $f_{\Omega_i, \Omega_v}(\Omega_i, \Omega_v) = dL(\Omega_i, \Omega_v)/dE(\Omega_v)$, where $\Omega_i$ is illumination direction and $\Omega_v$ is the viewing direction. Surface albedo results if the BRDF is integrated over the entire viewing hemisphere.

Vegetation indices emerge from the differences in reflectance at the red edge. However, BRDF emerges from geometric optics, that is, the shape and arrangement of objects within a pixel that transmit or block light (Torrance and Sparrow, 1967). When viewed from space, forests appear brighter as the view angle approaches the Sun angle and brightest when the observer is in line with the Sun, a position referred to as the hot spot (Fig. 1). This occurs because at zero

![Fig. 1](image_url) Canadian black spruce showing backscattering (sun behind observer) on the left and forward scattering (sun opposite observer) on the right. Note the bright region (hot spot) where all shadows are hidden on the left. The graphs show the classic drop-off of BRDF from the hot spot to the dark spot along the principle plane for needle-leaf forest. The red line shows the view zenith angle from each photo. Photograph by Don Deering.
phase angle, all shadows cast by objects are obscured (Hapke et al., 1996), and at higher phase angles more shadows become visible to the observer, which results in a darkening of the pixel’s reflectance. The rate of change of the reflectance with phase angle, in both the zenith and azimuth directions, is closely linked to the size, shape, and number of trees in the scene; and there is a quantitative theory of the shape of the BRDF (Li and Strahler, 1992). Observations of the BRDF from space demonstrate that less-structured scenes, such as deserts and tundra, have a flatter BRDF as a function of view zenith angle, and more-structured scenes such as savanna and forest have a more peaked BRDF (Béron et al., 1997; Bicheron and Leroy, 2000). In this study, we examine whether this technique could be used to detect the existence of tree-like structures on the surface of an extrasolar planet. As far as we are aware, this is the first study to examine the BRDF integrated over the entire planetary disk.

Spectral information from a Darwin- or TPF-type mission will be averaged over the entire disk, because the large distances to the planet prohibit greater spatial resolution, and averaged over a full rotation period, because spectral and temporal sampling will be limited by the relatively low brightness of the planet compared with its parent star. In this study, we developed a method to determine whether multicellular life, specifically canopy structure, could be detected on an Earth-like planet based on spectra that are averaged over the full disk for a full rotation period. We hypothesize that, as Earth approaches full illumination, an observer in line with the star and planet will observe an anomalous increase in albedo as shadows cast by vegetation are blocked by the vegetation (Fig. 2). Without vegetation there would be no anomalous increase in albedo as a planet approaches full illumination. We tested our theory by combining a well-tested BRDF model with a global vegetation map and global cloud data and simulations to model Earth’s albedo as viewed by a distant observer between phase angles of 0.5° and 44°.

2. Methods

To calculate disk and rotationally averaged planetary reflectance on a vegetated and non vegetated planet, we first modeled the reflectance for each grid cell at different geometries and phase angles (the angle between the star, the observer, and the planet), using the following semi-empirical model (Maignan et al., 2004; Bacour and Bréon, 2005), based on data from the POLDER instrument (Bicheron and Leroy, 2000):

\[ R(\Omega_0, \Omega_v, \varnothing) = k_0 + k_1 F_1(\Omega_0, \Omega_v, \varnothing) + k_2 F_2(\Omega_0, \Omega_v, \varnothing) \]

This equation calculates reflectance (R) for each vegetation type based on its geometry [solar zenith (\(\Omega_0\)), view zenith (\(\Omega_v\)), and relative azimuth (\(\varnothing\)) angles] between phase angles of 0.5° and 44°. It combines a geometric kernel (F1), which models a flat Lambertian surface covered with randomly distributed spheroids with the same optical properties as soil (Lucht et al., 2000), with a volumetric kernel (F2), which models a theoretical turbid vegetation canopy with high leaf density (Maignan et al., 2004). The model fixes the hot spot half-width parameter to 1.5°, the mean observed value (Bréon et al., 2002). K (\(k_0, k_1, k_2\)) are biome-dependant constants inverted from multi-angle reflectance data collected in the global POLDER satellite database (Bacour and Bréon, 2005). We calculated the reflectance for the visible (670 nm) and the near infrared (NIR) (865 nm) wavelengths. The choice of optimal wavelengths for this method is complicated and is explained in the discussion.

We calculated globally integrated planetary reflectance by combining a global vegetation map (2° by 2.5° resolution at the equator) (Oleson and Bonan, 2000) with surface reflectance values calculated for four plant functional types per pixel. The plant functional types used in the model are as follows: evergreen needle leaf forest, evergreen broadleaf forest, deciduous needle leaf forest, deciduous broadleaf forest, mixed forests, closed shrublands, open shrublands, woody savannas, savannas, grasslands, and croplands. Plant functional types distinguish growth form and hence BRDF properties. BRDFs for Earth could also be generated by using the MODIS global BRDF database; however, our method enables manipulation of the land surface.

Abundant multicellular life is more likely to arise on planets with liquid water, an oxygen atmosphere, and active geology (Kasting and Catling, 2003); and these will erode most steep geological features such as asteroid craters or most very steep mountains (>45° tilt). More than 99% of Earth’s surface has a slope less than 45° (Hall et al., 2005). We removed observations made at high phase angles (45°-90°) to

FIG. 2. A diagram of our proposed experimental design. The distant observer, a space telescope such as TPF, would observe the rate of change of albedo near the phase angles of 0.5° and 44°. \(\Omega\) is the azimuth angle, \(\Omega_0\) is the solar zenith angle, \(\Omega_v\) is the view angle, and \(\varnothing\) is the phase angle.
ensure any confounding effect from steep terrain is small. We also did not analyze phase angles between 45° and 90° to avoid forward scattering effects from water and clouds. There are only minor BRDF effects between phase angles of <45° for oceans (Jin et al., 2004), ice, and bare ground (Bicheron and Leroy, 2000). Sensitivity studies indicate that our results are still valid even if reflectance of oceans, ice, and bare ground increases by up to 0.06 at full illumination from backscattering BRDF effects.

We calculated planetary albedo as the integrated disk- and rotation-averaged emitted radiation over 144 simulations representing solar zenith 0° at the equator for each of the 144 longitudinal bands, each with separate view zenith, solar zenith, and azimuth angle for each pixel. In Fig. 3, we show view zenith, solar zenith, and azimuth angles for two such simulations, at 0.5° and 44° phase angles and their resulting reflectance. We assumed no tilt to the planet and further assumed that the observer is in an approximate plane with the planet’s orbit and its star. This assumption will be valid for Earth-sized planets detected by the Kepler observatory because it detects planets by using slight changes in the star’s brightness as a planet transits. The BRDF technique would require additional geometric calculations for planets that do not meet this assumption. By the cosine effect, pixels near the poles have proportionally less weighting. We used the following formula to calculate the area (A) of each grid cell on our maps:

\[ A = \int_{\varphi_1}^{\varphi_2} R \left( \int_{\lambda_1}^{\lambda_2} R' \cos \varphi d\lambda \right) d\varphi = R^2 (\lambda_2 - \lambda_1) (\sin \varphi_2 - \sin \varphi_1) \]

where longitudes are represented by \( \lambda_1 \) and \( \lambda_2 \), latitudes are represented by \( \varphi_1 \) and \( \varphi_2 \), with \( 0 \leq \lambda_1 \leq \lambda_2 \leq 2\pi \) and \( -\pi/2 \leq \varphi_1 \leq \varphi_2 \leq \pi/2 \). R is the Earth’s radius, and \( \lambda_1 \) and \( \lambda_2 \) are expressed in radians.

Reflected photon flux density of a planet will increase from dichotomy to full illumination as reflected light from a larger portion of the planet reaches the observer. Photons reaching the observer are a function of planetary albedo and the portion of the planet illuminated at different phase angles. In this paper, we define albedo as reflected photons divided by top of atmosphere incident radiation at full illumination.

We simulated global cloud cover using the National Center for Atmospheric Research Community Atmosphere Model (CAM 3.0; http://www.ccm.ucar.edu/models/atm-cam) (Collins et al., 2006), coupled with the Community Land Model (CLM 3.0; http://www.cgd.ucar.edu/tss/clm) and a slab ocean model. The simulations were run for 60 years at a 20-minute time step and a resolution of 2° by
2.5° at the equator. We simulated cloud cover for a vegetated Earth and a bare-ground Earth to see how vegetation can affect total planetary cloud cover. We combined simulated cloud height (low, medium, and high) and total percent cover with albedo values for low, medium, and high clouds (strato-cumulus, alto-stratus, and cirrus) at several planetary phase angles (Tinetti et al., 2006a). To calculate accurate monthly and annual cloud variability on Earth, we used 10 years (1986–1995) of an observational cloud data set, International Satellite Cloud Climatology.

Digitized data from Fig. 7 of Tinetti et al. (2006b) was used to obtain albedo for clouds (strato-cumulus, alto-stratus, and cirrus) at different planetary phase angles. Descriptions of the physics of this planetary model and further information on the data sets are available in papers by Tinetti et al. (2006a) and Tinetti et al. (2005). Albedo for each pixel was an area-weighted average of land surface albedo calculated by using our BRDF model multiplied by the percentage of the pixel not cloud covered and the albedo of low, medium, and high clouds based on height ratios and percentage cloud cover.

To test whether we could use this method to detect multicellular life under conditions of unknown planetary cloud cover, we varied the total planetary cloud cover between 50% and 70%. To understand how month-to-month changes in global cloud cover may impact albedo, we simulated albedo for each month separately at 50%, 60%, and 70% total planetary cloud cover. We then calculated the difference in the rate of change of albedo between phase angles 0.5° and 3.5° and phase angles 44° and 47° for all 36 simulations.

We tested our method based on two actual coronagraph designs (Guyon et al., 2006). The Achromatic Interferometric Coronagraph (Gay and Rabbia, 1996) has a 50% useful throughput (total amount of planet light, divided by the flux in planet light that is collected by the entrance aperture) at 0.5 λ d−1 (Guyon et al., 2006), which is close to the theoretical detection limit. λ is the wavelength of light; d is diameter of the aperture (m). This gives the minimum separation that can be detected between the star and the planet in radians, or the smallest inner working angle. If stellar angular size is considered, the best current coronagraph is the Phase Induced Amplitude Apodization Coronagraph (Guyon et al., 2005), which can detect planets up to 2 λ d−1. We calculated how close to its star we could detect an exoplanet based on the two coronagraph designs, a 12 m lens, 400 and 800 nm wavelengths, an Earth-like planetary radius of 1 AU, and varied distance to the star.

3. Results

A cloud-free planet with Earth-like vegetation increases its disk and rotationally averaged integrated albedo from 0.09 in the NIR (865 nm) and 0.06 in the visible (670 nm) at 44° phase angle to 0.18 in the NIR and 0.11 in the visible at 0.5° phase angle. Without vegetation, albedo increases from 0.12 in the NIR and 0.08 in the visible at a phase angle of 44° to 0.15 in the NIR and 0.11 in the visible at 0.5° phase angle. With cloud cover, the vegetated planet’s albedo increases from 0.17 in the NIR and 0.14 in the visible at 44° phase angle to 0.53 in the NIR and 0.38 in the visible at 0.5° phase angle. Without vegetation, a cloudy planet’s albedo increases from 0.18 in the NIR and 0.15 in the visible at 44° phase angle to 0.52 in the NIR and 0.38 in the visible at 0.5° phase angle. NIR albedo increases by 0.36 in the NIR and 0.24 in the visible between 0.5° and 44° on a vegetated planet with clouds and by 0.34 in the NIR and 0.23 in the visible on a non-vegetated planet with clouds, a difference of 0.02 and 0.01 (Table 1).

On Earth, clouds dominate planetary albedo, and large variations in cloud cover at different phase angles can cause an increase in brightness that mimics the BRDF effect of vegetation. Over a 10-year period, the yearly averaged International Satellite Cloud Climatology cloud cover averaged 0.64 ± 0.009. Planetary cloud cover can vary by ± 0.011 between the monthly maximum and the monthly minimum with a standard deviation of <1% (0.008) for all months. There were large regional differences between climate simulations for a vegetated planet and a non-vegetated planet, but month to month globally averaged planetary cloud cover only differed by 0.003 on average.

![FIG. 4. Planetary albedo at different phase angles at 865 nm for planets with 50% (dashed line), 60% (dotted line), 70% (solid line) cloud cover for a vegetated (gray) and non-vegetated (black) planet.](image-url)
Increasing cloud cover from 50% to 70% can increase albedo by \(0.12\) at 0.58 but only \(0.02\) at 448 phase angle (Fig. 4). Therefore, a planet with multicellular life cannot be distinguished by subtracting the albedo at phase angle 0.5° from albedo at phase angle 44° because this value will vary depending on total percentage cloud cover, which may not be known with accuracy.

Although an obstacle, it is possible to solve this by exploiting the differing rate of change of planetary albedo because the rate of increase of albedo is much greater between 0.5° and 3.5° than between 44° and 47° on a vegetated planet than on a non-vegetated planet (Fig. 5). Because of this differing rate of change, over a wide range of cloud cover, we were able to distinguish a vegetated planet from a non-vegetated planet, using the second derivative with respect to phase angle of planetary albedo as a planet rotates around its star (Fig. 6). We estimate that the rate of increase in brightness between 0.5° and 3.5° and 44° and 47° can distinguish multicellular life if planetary cloud cover is resolved to \(0.1\) in the NIR and \(0.05\) in the visible wavelengths (Fig. 6).

We used two coronagraph designs (2 and 0.5 λ d−1) at 400 nm (representing the visible spectrum) and 800 nm (representing the NIR spectrum) to estimate how close to its star a planet’s albedo can be measured. We estimate that a coronagraph 2 λ d−1 can measure a planet up to 2 light-years away within 1° of its star (Fig. 7). A coronagraph capable of 0.5 λ d−1 can measure a planet within 0.5° of its star up to 8 light-years away. At 800 nm, a coronagraph 0.5 λ d−1 can measure a planet within 0.5° of its star up to 4 light-years away, and a 2 λ d−1 coronagraph could measure up to 1 light-year away.

Based on estimates of how close to its star a planet can be resolved and how well a planet with multicellular life can be distinguished under different levels of planetary cloud cover, we estimate the theoretical maximum distance this technique could potentially be used to detect multicellular life. Using the rate of increase in brightness between 0.5° and 3.5° and 44° and 47° in the NIR, multicellular life may be detected if planetary cloud cover is known within ±0.1 (Fig. 6). A coronagraph 0.5 λ d−1 can detect an exoplanet within 0.5° of its star at 2 light-years (Fig. 7). Therefore, a 0.5 λ d−1 coronagraph at 800 nm wavelengths can potentially detect multicellular life only to 2 light-years if planetary cloud cover can be resolved to ±0.1 accuracy (Fig. 8). The optimal wavelength is 800 nm unless cloud cover can be resolved to better than ±0.03; then 400 nm is the optimal wavelength (Fig. 8). We assume that it will be difficult to resolve planetary cloud cover to better than ±0.02 and do not show this section of the graph because on Earth month-to-month cloud cover can vary by ±0.01.

### Table 1. The Increase in Albedo between Phase Angles 0.5° and 44° for a Simulated Planet with No Vegetation, Vegetation, Cloud Cover, and No Cloud Cover

<table>
<thead>
<tr>
<th></th>
<th>No vegetation</th>
<th>Vegetation</th>
<th>No vegetation + clouds ± sd</th>
<th>Vegetation + clouds ± sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Change in visible albedo</td>
<td>0.027</td>
<td>0.043</td>
<td>0.23 ± 0.007</td>
<td>0.24 ± 0.007</td>
</tr>
<tr>
<td>Change in NIR albedo</td>
<td>0.037</td>
<td>0.096</td>
<td>0.34 ± 0.012</td>
<td>0.36 ± 0.011</td>
</tr>
</tbody>
</table>

Cloud cover is the average planetary cloud cover for each of the 12 months \(N = 12\). sd, standard deviation.

### 4. Discussion

A planet with multicellular life would have a peak in brightness as it approaches full illumination due to anisotropic effects that would not be present on a lifeless planet.

**FIG. 5.** The difference in the rate of change of albedo at 865 nm between phase angles 0.5° and 3.5° and 44° and 47° for a vegetated planet with no clouds (top), a cloudy planet with no vegetation (middle), and a cloudy planet with vegetation (bottom).
with abundant water or a planet covered with microbial life. Assuming that most physical processes that affect cloud formation on extrasolar planets habitable to multicellular life are similar to those on Earth, variations in cloud cover or height should not change enough between dichotomy and full illumination to mimic a vegetation hot spot. However, we cannot just subtract the albedo at phase angle $0.5^\circ$ from albedo at phase angle $44^\circ$ to determine whether there is vegetated life on a planet because this number will vary depending on total percentage cloud cover. If the rotation period can be determined with accuracy on an exoplanet, deviations from a periodic signal can be used to infer the

![FIG. 6. Change in the rate of increase of integrated planetary albedo in the NIR (top) and visible wavelengths (bottom) for a vegetated planet (gray) and a planet with no vegetation (black) between phase angles $0.5^\circ$ and $3.5^\circ$ and $44^\circ$ and $47^\circ$ at different levels of planetary cloudiness (50–70%). Error bars (standard deviation) show differences between 12 simulations representing average global cloud cover for each month of the year.](image)

![FIG. 7. The minimum distance an exoplanet can be resolved from its star (degrees) using two potential coronagraph designs, $2 \lambda d^{-1}$ (dashed lines) and $0.5 \lambda d^{-1}$ (solid lines), and two wavelengths, 400 nm (gray) and 800 nm (black), assuming an Earth-like orbit (1 AU).](image)
presence of relatively short-lived structures such as clouds (Palle et al., 2008).

Even if cloud cover cannot be determined with accuracy, we can exploit the differing rate of change of planetary albedo between vegetated and non-vegetated planets. This capitalizes on the fact that clouds and trees increase in brightness at distinctly different rates as a planet approaches full illumination: clouds exhibit forward scattering (Mishchenko et al., 1996; Gatebe et al., 2003), while trees exhibit backscattering (Bicheron and Leroy, 2000). This means that trees will increase in brightness as a planet approaches full illumination, as the shadows are hidden; but clouds will increase much less.

According to our simulations, we can detect multicellular life on an Earth-like planet by using the 2nd derivative of planetary albedo with respect to phase angle between phase angles 0.5 and 3.5 and 44 and 47. Planetary cloud cover can vary by ±0.01 between the monthly maximum and the monthly minimum. However, this month-to-month change in cloud cover has only minor effects on the 2nd derivative with respect to phase angle of brightness as shown by the small error bars in Fig. 6. This additional constraining observation of BRDF may be sufficient to discriminate multicellular life from single-cellled life on extrasolar planets. In addition, using the rate of change of albedo will eliminate the need to estimate ground albedo accurately, which is not likely to be known with accuracy.

A dry lifeless planet similar to the Moon exhibits a strong hot spot due to shadows from craters and very dry fine soils that cause coherent backscatter. Coherent backscatter can occur on very dry soils where particles have a diameter that is similar to the wavelength of the photon used to view them. This hot spot occurs because portions of wave fronts that are multiply scattered within a non-uniform, wavelength-sized medium combine after following the same path, but in opposite directions (Hapke et al., 1993). In this study, we assumed that there would be no coherent backscatter on wet planets capable of sustaining multicellular life when the planets are viewed with wavelengths of 800 nm or less. On Earth, dry areas, such as deserts, generally have particle sizes of 0.05 to 2 mm (Tarbuck et al., 2008), which are too big to exhibit coherent backscatter at 800 nm wavelengths. Desert areas can also exhibit hot spot features through geometric shadow processes, but the amplitude of the change in reflectance between the hot spot and dark spot is small, as exemplified by the BRDF of a barren region of Libya (Bicheron and Leroy, 2000). Wet areas would not exhibit coherent backscatter because extremely dry conditions (such as those present on the Moon) are necessary (Hapke et al., 1996).

Mountains, asteroid craters, or other shadow-casting geological features can potentially confound the attribution of BRDF to multicellular life. However, most geological features such as asteroid craters or steep mountains (>45° slope) will be eroded by abundant liquid water, an oxygen atmosphere, and an active geology, all of which are likely to be present on any planet capable of sustaining multicellular life (Kasting and Catling, 2003). A lifeless planet with a similar climate to Earth will be topologically very similar to Earth, although most sediment will be less weathered (Dietrich and Perron, 2006). Therefore, we can assume that, in terms of slope, Earth is typical of wet terrestrial planets with <1% of the surface with a slope greater than 45° (Hall et al., 2005). We removed observations made at phase angles larger than 45° to ensure that any confounding effects of

![Graph](image_url)
steep terrain are small. On other planets, as on Earth, there may be unique topological features, like the hoodoo formations in Bryce Canyon or the limestone formations in Guilin, China, that exhibit backscattering anisotropic effects. However, hoodoo structures will likely be uncommon (<1% for the surface), as they need a rare combination of basalt and tuff to form and will eventually succumb to erosion. Limestone is of mainly marine biogeochemical origin, from shells and corals (Tarbuck et al., 2008), and is likely to be less abundant on a lifeless planet.

The ocean and clouds dominate the albedo of Earth and will play an important role in determining the albedo of any exoplanet with abundant liquid water. The ocean and clouds forward scatter light, while trees backscatter light. Most of the change in reflectance from anisotropic effects for clouds (Mishchenko et al., 1996) and the ocean (Jin et al., 2004) is from phase angles >45°. By removing phase angles >45°, we can also remove confounding anisotropic effects of clouds and oceans. Typical ocean glint is 0.2 in the specular direction (Bréon and Henriot, 2006). However, reflectance in the specular direction will only rarely replicate the effect of a vegetation hot spot and, therefore, will not impact our results.

Is this method as useful for detecting life on exoplanets as using the red edge? The change in reflectance between the hot spot and the dark spot of a forest can be as large as the change in reflectance between the visible and NIR of a forest. For instance, the reflectance of a tropical broadleaf evergreen forest can increase by 0.3 between the hot spot and the dark spot (Bicheron and Leroy, 2000). This is similar to the increase in surface reflectance of a forest from 0.03 at 400 nm to 0.27 at 800 nm (Tinetti et al., 2006a). Previous simulations have shown the difficulty of resolving the red edge if an exoplanet is resolved to a single pixel (Tinetti et al., 2006b). Our simulations suggest that, by using advanced coronagraph designs and accurately resolving planetary cloud cover, a BRDF signal of life could be as large as a red-edge signal for nearby stellar systems.

To determine the optimal wavelength for this technique, it is first necessary to know how well cloud cover on an exoplanet can be resolved (Palle et al., 2008). We determine that, if planetary cloud cover can be resolved to better ~±0.03, then 400 nm wavelength is optimal; otherwise, 800 nm wavelength is optimal (Fig. 8). In theory, a broadband albedo would work as well as a narrowband albedo. It has been hypothesized that planets with dimmer stars may have a red edge at higher wavelengths (Wolstencroft and Raven, 2002). In such a case, additional calculations will be necessary to determine optimal wavelengths.

How well this method could work is dependent on coronagraph design (Guyon et al., 2006) and how well cloud cover can be resolved (Palle et al., 2008) (Fig. 8). There are ~50 stellar systems within 5 parsecs of Earth. If cloud cover on exoplanets can be resolved to better than ±0.03, then we estimate that this method may be effective in some of these stellar systems with currently possible coronagraph designs (2 λ d−1) and on all of these stellar systems if future coronagraphs can be designed to resolve exoplanets to within 0.5 λ d−1 of their star (Table 2). Recently, an exoplanet 129 light-years away was resolved at 2 λ d−1 with a small (1.5 m) ground-based telescope (Serabyn et al., 2010). This demonstrates the potential of this method to be used with ground-based telescopes capable of much larger lens diameters that could further expand the range of this technique.

We can envision a process where a terrestrial exoplanet’s size, density, cloud cover, distance to star, and the star’s irradiance are measured or estimated. Based on these data, a model can be developed with which to estimate how albedo would change between 0° and 45° phase angles, assuming a relatively smooth, eroded, planetary surface. These simulations can be compared to the actual rate of change of brightness observed. If the rate of change of brightness between 0.5° and 3° and 44° and 47° is greater than the modeled estimate, then this is evidence for multicellular life on the planet. False positives caused by instrument error or intermittent events such as volcanic activity or changing cloud cover could be determined by observing the planet during continuous rotation cycles. Multicellular life would continuously demonstrate the BRDF signal, while other causes would demonstrate it only intermittently.

| 5. Conclusions |

We can articulate four archetypical scenarios of planetary life: a dry lifeless planet, a lifeless planet with abundant liquid water, a planet with abundant liquid water and microbial life, and a planet with abundant liquid water and multicellular life (Table 3). Ideally, a set of observational constraints can be made that can distinguish these characteristics for a candidate planet. A dry lifeless planet will likely exhibit a hot spot due to abundant craters and wavelength-sized dust (Hapke et al., 1993), but it will not exhibit signs of clouds or atmospheric water vapor. A lifeless planet with abundant liquid water will likely not exhibit a hot spot but will exhibit signs of clouds and atmospheric water. A planet with abundant liquid water and single-cellled life will exhibit clouds, water, potentially a red-edge signal, but no BRDF effect (Tinetti et al., 2006b). A planet with abundant liquid water and abundant multicellular life will exhibit signs of clouds, water, red edge, and a BRDF effect. If anisotropic effects are taken into account, tree-like multicellular life may be distinguished from single-cellled life on an exoplanet.

| Table 2. The Distance to an Exoplanet That Multicellular Life May Be Detected, Assuming a 12-Meter Diameter Aperture Telescope and the Optimal Wavelength (400 or 800 nm) Based on Cloud Resolution Accuracy (±) and Coronagraph Design |
|-----------------|-----------------|-----------------|-----------------|
| 2 λ d−1          | 1 light-year     | 4 light-years   | 6 light-years   |
| 0.5 λ d−1        | 3 light-years    | 14 light-years  | 18 light-years  |

| Table 3. Four Scenarios of Planets with Distinct Spectra for Indicators of Life |
|-----------------|-----------------|-----------------|-----------------|
| Dry lifeless    | No              | No              | Yes             |
| Lifeless + water| Yes             | No              | No              |
| Microbial + water| Yes            | Maybe           | No              |
| Multicellular + water| Yes   | Yes             | Yes             |
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Abbreviations

BRDF, bidirectional reflectance distribution function; NIR, near infrared; TPF, Terrestrial Planet Finder.

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