11

12

13

15

16

17

18

19

20

21

22

23

25

26

27

29

30

31

32

Bioverse: Potentially Observable Exoplanet Biosignature Patterns Under the UV Threshold Hypothesis for the Origin of Life

Martin Schlecker , 1 Dániel Apai , 1, 2 Antonin Affholder , 3 Sukrit Ranjan , 2, 4 Régis Ferrière , 3, 5, 6 Kevin K. Hardegree-Ullman , 1, 7 Tim Lichtenberg , 8 and Stéphane Mazevet , 9

1 Steward Observatory, The University of Arizona, Tucson, AZ 85721, USA; schlecker@arizona.edu

2 Lunar and Planetary Laboratory, The University of Arizona, Tucson, AZ 85721, USA

3 Department of Ecology and Evolutionary Biology, University of Arizona, Tucson AZ, USA

4 Blue Marble Space Institute of Science, Seattle, 98104, USA

5 Institut de Biologie de l'École Normale Supérieure, ENS, PSL, Paris, France

6 International Research Laboratory for Interdisciplinary Global Environmental Studies (iGLOBES), CNRS, ENS, PSL, University of Arizona, Tucson AZ, USA

7 Caltech/IPAC-NASA Exoplanet Science Institute, 1200 E. California Blvd., MC 100-22, Pasadena, CA 91125, USA

8 Kapteyn Astronomical Institute, University of Groningen, PO Box 800, 9700 AV Groningen, The Netherlands

9 Observatoire de la Côte d'Azur, Université Côte d'Azur, Nice, France

ABSTRACT

A wide variety of scenarios for the origin of life have been proposed, with many influencing the prevalence and distribution of biosignatures across exoplanet populations. This relationship suggests these scenarios can be tested by predicting biosignature distributions and comparing them with empirical data. Here, we demonstrate this approach by focusing on the cyanosulfidic origins-of-life scenario and investigating the hypothesis that a minimum near-ultraviolet (NUV) flux is necessary for abiogenesis. Using Bayesian modeling and the Bioverse survey simulator, we constrain the probability of obtaining strong evidence for or against this "UV Threshold Hypothesis" with future biosignature surveys. Our results indicate that a correlation between past NUV flux and current biosignature occurrence is testable for sample sizes of ≥ 50 planets. The diagnostic power of such tests is critically sensitive to the intrinsic abiogenesis rate and host star properties, particularly maximum past NUV fluxes. Surveys targeting a wide range of fluxes, and planets orbiting M dwarfs enhance the chances of conclusive results, with sample sizes \$\ge 100\$ providing \$\ge 80\% likelihood of strong evidence if abiogenesis rates are high and the required NUV fluxes are moderate. For required fluxes exceeding a few hundred erg/s/cm², both the fraction of inhabited planets and the diagnostic power sharply decrease. Our findings demonstrate the potential of exoplanet surveys to test origins-of-life hypotheses. Beyond specific scenarios, this work underscores the broader value of realistic survey simulations for future observatories (e.g., HWO, LIFE, ELTs, Nautilus) in identifying testable science questions, optimizing mission strategies, and advancing theoretical and experimental studies of abiogenesis.

1. INTRODUCTION

The probability of abiogenesis remains one of the fundamental questions in modern science. As of now, it has been primarily explored through statistical arguments (e.g., Spiegel & Turner 2012; Kipping 2021; Lingam et al. 2024). At the same time, a wide variety of scenarios for the origin of life have been proposed (e.g., Baross & Hoffman 1985; Brasier et al. 2011; Mulkidjanian et al. 2012; Fox & Strasdeit 2013; Deamer & Georgiou 2015; Westall et al. 2018). While we may still be far from conclusively testing these scenarios, new prospects in the search for conditions favorable to life have opened up by thinking

of the origin of life as a planetary phenomenon and identifying global-scale environmental properties that might support pathways to life (Sasselov et al. 2020). In particular, specific planetary conditions are needed to create stockpiles of initial compounds for prebiotic chemistry; and planetary processes are required to trigger the prebiotic synthesis. Such planetary conditions can be hypothesized for exoplanets located in the habitable zone (HZ) of their host star, with persistent liquid water on their surface. For example, if deep-sea or sedimentary hydrothermalism is required for abiogenesis, then the insulation of an ocean from the planetary crust minerals (e.g., due to high-pressure ices) may reduce or eliminate the chances of life emerging (e.g., Baross & Hoffman 1985). The alternate scenario of a surface locally subject to wet-dry cycles requires a planetary exposure to mid-range ultraviolet (UV) irradiation, as a source of energy and an agent of selection in chemical evolution (e.g., Deamer et al. 2019). This "UV Threshold Hypothesis" states that UV light in a specific wavelength range played a constructive role in getting life started on Earth (Ranjan & Sasselov 2016; Ranjan et al. 2017a; Rimmer et al. 2018; Rapf & Vaida 2016), and it could provide a probabilistic approach to the interpretation of possible future biosignature detections (e.g., Catling et al. 2018a; Walker et al. 2018).

The association of chemical pathways to life and plan-72 etary environmental conditions offers a new opportunity 73 to test alternate scenarios for life emergence based on 74 planetary-level data collected from the upcoming ob-75 servations of populations of exoplanets. Deep-sea hy-76 drothermal scenarios require planetary conditions that 77 may not be met on ocean worlds with large amounts 78 of water, where the water pressure on the ocean floor 79 is high enough to form high-pressure ices (Noack et al. 80 2016; Kite & Ford 2018). In this case, a testable predic-1 tion would be that planets with high-pressure ices do not show biosignatures. Likewise, if UV light is required to 83 get life started, then there is a minimum planetary UV 84 flux requirement to have an inhabited world. This re-85 quirement is set by competing thermal processes; if the 86 photoreaction does not move forward at a rate faster 87 than the competitor thermal process(es), then the abio-88 genesis scenario cannot function. On the other hand, 89 abundant UV light vastly in excess of this threshold 90 does not increase the probability of abiogenesis, since once the UV photochemistry is no longer limiting, some 92 other thermal process in the reaction network will be 93 the rate-limiting process instead. Therefore, a putative dependence of life on UV light is best described as a step 95 function (e.g., Ranjan et al. 2017a; Rimmer et al. 2018, 96 2021a).

The goal of this work is to evaluate the potensial of future exoplanet surveys to test the hypothesis that a minimum past near-ultraviolet (NUV) flux is required for abiogenesis. We focus on one version of the UV Threshold Hypothesis, the so-called cyanosulfidic scenario, which has been refined to the sured to be $(6.8 \pm 3.6) \times 10^{10}$ photons cm⁻² s⁻¹ nm⁻¹ integrated from 200–280 nm at the surface (Rimmer test al. 2018, 2021b; Rimmer 2023; Ranjan et al. 2023)¹.

While prebiotic photochemistry is fundamentally driven by photon number flux, we choose
to frame the problem in terms of energy flux,
for which empirical estimates in the NUV extst across stellar populations and ages (RicheyYowell et al. 2023). Due to the inherent uncertainties concerning the relationship between surface and top-of-atmosphere fluxes, we treat the
threshold NUV flux as a free parameter.

We first follow a semi-analytical Bayesian analysis to estimate probabilities of obtaining strong evidence for or against the UV Threshold Hypothesis. Under this hypothesis (H_1) , the probability of an exoplanet having detectable biosignatures is zero if the NUV irradiation is less than the threshold, and it is equal to the (unknown) probability of life emerging and persisting, f_{life} , if NUV exceeds the threshold for a sufficiently long period of time. Under the null hypothesis (H_{null}) , that probability simply is f_{life} , that is, it does not correlate with the UV flux.

Figure 1 shows these hypotheses as derived from the predictions of the cyanosulfidic scenario. Given a sample of planets, where for some of them we have convincing biosignature detections but remain agnostic on $f_{\rm life}$, we have evidence for H_1 and $H_{\rm null}$ we can expect to obtain.

A real exoplanet survey will be subject to observational biases and sample selection effects, and will
be constrained by the underlying demographics of the
planet sample. To assess the information gain of a realistic exoplanet survey, we employed Bioverse (Bixel
Apai 2021; Hardegree-Ullman et al. 2023; Schlecker
et al. 2024; Hardegree-Ullman et al. 2025), a framework that integrates multiple components including statistically realistic simulations of exoplanet populations,
a survey simulation module, and a hypothesis testing
module to evaluate the statistical power of different ob-

This paper is organized as follows: In Section 2, we introduce both our semi-analytical approach and Bioverse simulations for testing the UV Threshold Hyments pothesis. Section 3 presents the results of these experiments for a generic survey as well as for a realistic transition sit survey. In Section 4, we discuss our findings before concluding with a summary in Section 5.

2. METHODS

152

153

154

2.1. Fraction of inhabited planets with detectable biosignatures

Here, we conduct a theoretical experiment on the UV Threshold Hypothesis by relating the occurrence of life on an exo-earth candidate with a minimum past

 $^{^{1}}$ or $\sim 43\pm 23\,\mathrm{erg\,s^{-1}\,cm^{-2}}$ in energy flux units.

Hypothesis: Life only originates on planets with particular UV irradiance

Prediction

H1 Correlation between past UV flux and biosignature occurrence

Past UV flux and the occurrence of biosignatures are correlated.

164

190

H0 No correlation between past UV flux and biosignature occurrence

Figure 1. UV Threshold Hypothesis and null hypothesis derived from the cyanosulfidic scenario.

158 quiescent stellar UV flux, focusing on the prebiotically 159 interesting NUV range from 200–280 nm (Ranjan & Sas-160 selov 2016). Our core hypothesis shall be that life only 161 occurs on planets that at some point in their history have 162 received such radiation at a flux exceeding a threshold 163 $F_{\rm NUV,min}$.

2.2. Semi-analytical approach

We first assessed the expected probabilities of obtaining true negative or true positive evidence for the UV Threshold Hypothesis (H_1) above, as well as the probability for misleading or inconclusive evidence, under idealized conditions. This serves as a first-order estimate of the information content of a survey, before we take into account the effects of exoplanet demographics, sample selection, and survey strategy.

Presumably, not all habitable worlds are inhabited and not all inhabited worlds develop detectable biosignatures. The fraction of exo-Earth candidates (EEC) that 176 are both inhabited and exhibit detectable biosignatures the time of observation is unknown and is repre-178 sented by the term f_{life} . This encompasses the 179 probability of life both emerging and persisting 180 to produce detectable biosignatures. Due to our 181 ignorance about its true value or even its order of 182 magnitude, we draw $f_{
m life}$ from a log-uniform prior 183 probability distribution (but see Appendix B for 184 a discussion of the impact of different priors). Let us consider the probability to detect a biosignature 186 P(L), and let our observable be the inferred past NUV 187 flux of the planet F_{NUV} . Under Hypothesis H_1 , there exists a special unknown value of F_{NUV} , noted $F_{\text{NUV,min}}$ 189 such that

$$P(L|F_{\text{NUV}}, H_1) = f_{\text{life}} \quad \text{if } F_{\text{NUV}} > F_{\text{NUV,min}}$$
 (1)

$$P(L|F_{\text{NUV}}, H_1) = 0$$
 otherwise (2)

where $f_{\rm life}$ is the unknown probability of abiogenesis. The corresponding null hypothesis $H_{\rm null}$ is that there exists no such special value of $F_{\rm NUV}$ and that

$$P(L|F_{\text{NUV}}, H_{\text{null}}) = f_{\text{life}}.$$
 (3)

¹⁹⁷ In other words, H_{null} states that P(L) is independent of F_{NUV} .

Next, we determine the probability distribuwe tion of sample outcomes, or likelihood of each hypothesis. Let $Y = \sum_{i=1}^{n} L_i$ be the random variable counting the number of positive life detections in a sample of size n. Its probability mass function under the null hypothesis $H_{\rm null}$ is that of a binomial distribution:

$$P(Y = k|H_{\text{null}}) = \binom{n}{k} f_{\text{life}}^k (1 - f_{\text{life}})^{n-k}. \tag{4}$$

Under H_1 , Y also follows a binomial distribution, however it is conditioned by $n_{\lambda}=n(\{F_{\mathrm{NUV,i}} \text{ if } F_{\mathrm{NUV,i}} > 208\ F_{\mathrm{NUV,min}}\}_{i\in[1,n]})$, the number of values of F_{NUV} in the experiment that exceed $F_{\mathrm{NUV,min}}$

$$P(Y = k|H_1) = \binom{n_{\lambda}}{k} f_{\text{life}}^k (1 - f_{\text{life}})^{n_{\lambda} - k}.$$
 (5)

Following a similar approach as in Affholder et al. (2025), we aim to quantify the information gain from our sampling procedure by computing the Bayes factors (Jeffreys 1939)

$$BF_{H_1,H_{\text{null}}} = \frac{P(Y = k|H_1)}{P(Y = k|H_{\text{null}})} = \frac{\binom{n_{\lambda}}{k}}{\binom{n}{k}} (1 - f_{\text{life}})^{n_{\lambda} - n}$$
215 (6)

216 and

$$BF_{H_{\text{null}},H_1} = \frac{P(Y=k|H_{\text{null}})}{P(Y=k|H_1)} = \frac{\binom{n}{k}}{\binom{n_{\lambda}}{k}} (1 - f_{\text{life}})^{n-n_{\lambda}}.$$

²¹⁸ Given a sample of planets, where for some of them we have convincing biosignature detections but remaining agnostic on $f_{\rm life}$: What evidence for H_1 and $H_{\rm null}$ can we expect to get? The analytical expression for the Bayes factor of this inference problem (Equation 6) is determined by the unknown variables $f_{\rm life}$ and $F_{\rm NUV,min}$, as well as by the summary statistic Y (number of biosignature detections). To compute the distribution of evidences, we repeatedly generated samples under H_1 and $H_{\rm null}$ and computed the Bayes factors H_1 and H_1 and H_2 we then evaluated the fraction of Monte

237

240

241

243

244

245

247

248

249

250

251

252

254

255

256

257

258

259

261

262

264

265

266

267

268

269

270

271

272

273

275

229 Carlo runs in which certain evidence thresholds (Jeffreys 1939) were exceeded.

Under a more realistic scenario, the distribution of 232 n_{λ} depends on additional planetary properties and their evolution, as well as on observational biases and sample 234 selection effects of the survey. We will address these in 235 the following section.

2.3. Exoplanet survey simulations with Bioverse

To assess the diagnostic power of realistic exoplanet 238 surveys, we employed our survey simulator and hypothesis testing framework Bioverse (Bixel & Apai 2021). The general approach is as follows:

- 1. Exoplanet population synthesis: We populate the Gaia Catalogue of Nearby Stars (Smart et al. 2021) with synthetic exoplanets whose orbital parameters and planetary properties reflect our current understanding of exoplanet demographics (Bergsten et al. 2022; Hardegree-Ullman et al. 2023). Here, we also inject the demographic trend in question - in this case we assign biosignatures according to H_1 , i.e., to planets in the HZ that have received NUV fluxes above a certain threshold.
- 2. Survey simulation: We simulate the detection and characterization of these exoplanets with a hypothetical survey, taking into account the survey's sensitivity, target selection, and observational biases. To model the sensitivity of the information gain of a proposed mission to sample selection and survey strategy, we conduct survey simulations with Bioverse using different sample sizes and survey strategies.
- 3. **Hypothesis testing:** We evaluate the likelihood that a given survey would detect a specified demographic trend in the exoplanet population and estimate the precision with which the survey could constrain the parameters of that trend. A common definition of the null hypothesis H_{null} , which is also applied here, is that there is no relationship between the independent variable (here: maximum NUV flux) and the dependent variable (here: biosignature occurrence). The alternative hypothesis H_1 proposes a specific relationship between the independent and dependent variables. Bioverse offers either Bayesian model comparison or non-parametric tests to evaluate the evidence for or against the null hypothesis.

To determine the diagnostic capability of a given sur-276 vey, Bioverse runs multiple iterations of the simulated 278 survey and calculates the fraction of realizations that 279 successfully reject the null hypothesis. We used this 280 metric, known as the statistical power, to quantify the 281 potential information content of the survey, identify crit-282 ical design trades, and find strategies that maximize the 283 survey's scientific return.

2.3.1. Simulated star and planet sample

We generated two sets of synthetic exoplanet popula-286 tions, one for FGK-type stars and one for M-type stars. 287 The stellar samples are drawn from the Gaia Catalogue 288 of Nearby Stars (Smart et al. 2021) with a maximum 289 Gaia magnitude of 16 and a maximum stellar mass of 290 1.5 M_{\odot} . We included stars out to a maximum distance d_{max} that depends on the required planet sample size. 292 Planets were generated and assigned to the synthetic 293 stars following the occurrence rates and size/orbit distributions of Bergsten et al. (2022). Following Bixel & 295 Apai (2021), we considered only transiting EECs with 296 radii $0.8 S^{0.25} < R < 1.4$ that are within the HZ (see Section 2.3.2). The lower limit was suggested as a min-298 imum planet size to retain an atmosphere (Zahnle & ²⁹⁹ Catling 2017). For all survey simulations and hypothe-300 sis tests, we repeated the above in a Monte Carlo fashion 301 to generate randomized ensembles of synthetic star and 302 planet populations (Bixel & Apai 2021).

2.3.2. Habitable zone occupancy and UV flux

303

To construct a test of the UV Threshold Hypothe-305 sis, we required that life occurs only on planets with 306 sufficient past UV irradiation exceeding the origins of 307 life threshold $F_{\rm NUV,min}$. Further, we required this flux 308 to have lasted for a minimum duration $\Delta T_{\rm min}$ to al-309 low for a sufficient "origins timescale" (Rimmer 2023). 310 We nominally adopt $\Delta T_{\min} = 1 \,\text{Myr.}$ Under H_1 , 311 longer origins timescales have minimal impact 312 on a generic transit survey but significantly de-313 crease the fraction of inhabited planets around 314 FGK stars, as explored in Appendix C.

All commonly investigated origins-of-life scenarios re-316 quire water as a solvent; we thus considered only rocky 317 planets that may sustain liquid water on their surface, 318 i.e., that occupy their host star's momentary HZ during the above period, as well as at the epoch of observation. 320 To determine HZ occupancy, we took into account the 321 evolution of the host star's luminosity and HZ bound-322 aries.

The HZ describes a region around a star where a 324 planet with Earth's atmospheric composition and cli-325 mate feedbacks can maintain liquid water on its surface (e.g., Ramirez & Kaltenegger 2017; Ramirez 2018; Mol Lous et al. 2022; Spinelli et al. 2023; Tuchow &

Wright 2023). Here, we adopted orbital distance estimates that define the HZ as the region between the run-330 away greenhouse transition, where the stellar instella-331 tion cannot anymore be balanced through infrared cooling to space (Ingersoll 1969), and the maximum green-333 house limit, corresponding to the maximum distance at 334 which surface temperatures allowing liquid water can be maintained through a CO₂ greenhouse effect (Kast-336 ing 1991; Kasting et al. 1993; Underwood et al. 2003; 337 Kopparapu et al. 2013, 2014). The exact bound-338 aries of the habitable zone are known to be 339 sensitive to the star's luminosity, spectral type, 340 the planet's mass, and the planet's atmospheric properties (e.g., Pierrehumbert & Gaidos 2011; 342 Ramirez & Kaltenegger 2014, 2017, 2018; Koll 343 & Cronin 2019; Ramirez 2018, 2020; Bonati & Ramirez 2021; Chaverot et al. 2022; Turbet et al. 345 2023). Here, we adopted the commonly used 346 parametrization in Kopparapu et al. (2014) to 347 derive luminosity and planetary mass-dependent distance limits of the HZ $a_{\rm inner}$ and $a_{\rm outer}$.

To determine HZ occupancy, we interpolated the stellar luminosity evolution grid of Baraffe et al. (1998) using a Clough Tocher interpolant (Nielson 1983; Alfeld
1984, see left panel of Figure 2) to compute the evoluindicated the inner (runaway greenhouse) and outer (maximum greenhouse) edges as a function of planet mass
and stellar spectral type (Kopparapu et al. 2014). Being a local interpolation method, Clough Tocher enables
rapid processing while producing a smooth interpolating
surface that highlights local trends. From this, we get
each planet's epochs within and outside the HZ.

For the NUV flux, we used the age- and stellar mass- dependent NUV fluxes in the HZ obtained by Richey-Yowell et al. (2023), which considers GALEX UV data in the wavelength range of 177–283 nm. We linearly interpolate in their measured grid, where we convert spectral type to stellar mass using the midpoints of their mass ranges (0.75 $\rm M_{\odot}$ for K stars, 0.475 $\rm M_{\odot}$ for early-type M stars, and 0.215 $\rm M_{\odot}$ for late-type M stars). Outside the age and stellar mass range covered in Richey-Yowell et al. (2023), we extrapolate using nearest simplex (see right panel of Figure 2).

We then determined which planets were both in the HZ and had NUV fluxes above $F_{\rm NUV,min}$. To avoid considering short transitional phases, we require this situation to last for a minimum duration $\Delta T_{\rm min} \geq 1\,{\rm Myr}$. We assigned the emergence and persistence of life to a random fraction $f_{\rm life}$ of all temperate planets fulfilling these requirements. For the probability of a planet having detectable biosignatures, $P({\rm bio})$, the UV Threshold

379 Hypothesis then states

380
$$H_1: P(\text{bio}) = \begin{cases} 0, & F_{\text{NUV}} < F_{\text{NUV,min}} \\ f_{\text{life}}, & F_{\text{NUV}} \ge F_{\text{NUV,min}} \\ & \text{and in HZ for } \Delta t \ge 1 \text{ Myr} \end{cases}$$
 (8)

381 and the corresponding null hypothesis $H_{\text{null}}: P(\text{bio}) =$ 382 f_{life} , i.e., no correlation with UV flux.

2.3.3. Transit survey simulations

With the synthetic star and planet samples gener-384 ated, we used Bioverse's survey module to simulate 386 noisy measurements of key observables with a transit 387 survey. We assumed a hypothetical mission that can 388 target a large planet sample with high photometric pre-389 cision and conduct a biosignature search on these planets (e.g., Apai et al. 2019, 2022). The simulated survey 391 was designed to measure planetary instellation (for HZ 392 occupancy) with a precision of 5% and host star effec-393 tive temperature with a precision of 50.0 K. The maximum past NUV flux a planet received can be determined within a precision of 5%. To marginalize over choices of 396 biosignatures and their detectability, which are beyond 397 the scope of this study, we assumed that any inhabited 398 planet would show a biosignature detectable by the surз99 vey.

2.3.4. Hypothesis testing

We ought to choose a statistical test that is sen-401 402 sitive to the UV Threshold Hypothesis, and that 403 could be realistically conducted in a future tran-404 sit survey (which may include auxiliary informa-405 tion from ground-based observations, archived 406 data, or models). Given the available types 407 of data expected from such surveys, our test 408 shall be non-parametric and compare two sam-409 ples - planets with and without biosignatures $_{410}$ – to assess whether they are drawn from the 411 same underlying population in terms of their in-412 fered historic maximum NUV flux. Common op-413 tions include the Kolmogorov-Smirnov test, the 414 Brunner-Munzel test (Brunner & Munzel 2000), 415 and the Mann-Whitney U test (Mann & Whit-416 ney 1947). Due to its availability and suitabil-417 ity for large sample sizes, we chose the Mann-418 Whitney U test, which evaluates if one sample 419 is stochastically greater than the other. Here, 420 we compare the distributions of NUV fluxes of planets 421 with and without biosignatures. The implementation 422 in Bioverse relies on the scipy.stats.mannwhitneyu function (Virtanen et al. 2020) and returns a p-value for 424 each test. To balance the trade-off between Type

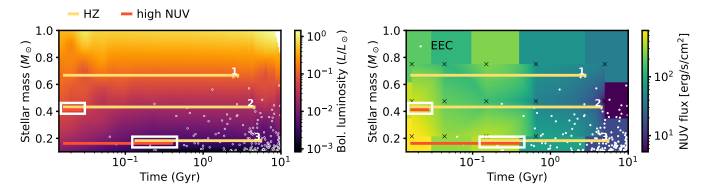


Figure 2. Interpolated stellar luminosity evolution (left) and evolution of the NUV flux in the HZ (right) as a function of host star mass. Scatter points show age and host star mass of the transiting planets in the synthetic planet sample; crosses denote the estimated NUV values in Richey-Yowell et al. (2023). We show three evolutionary tracks for a threshold flux of $F_{\text{NUV,min}} = 300.0 \,\text{erg}\,\text{s}^{-1}\,\text{cm}^{-2}$ that occupy the HZ (yellow sections) and exceed the threshold NUV flux (red sections) at different times. Where these sections overlap (white rectangles), the requirements for abiogenesis are met and we assign a biosignature detection with probability f_{life} . Planet 1 is an EEC orbiting a K dwarf that never receives sufficient NUV flux for abiogenesis. Planet 2 and Planet 3 enter the HZ at different times and receive sufficient NUV flux for different durations until their respective host star evolves below the threshold.

483

 $_{425}$ I and Type II error risks, we set the significance $_{426}$ level to the widely adopted threshold $\alpha=0.05$. $_{427}$ To quantify the diagnostic power of the survey, $_{428}$ we conducted repeated randomized realizations $_{429}$ and calculated the fraction of successful rejections of the null hypothesis, i.e., the statistical $_{431}$ power.

3. RESULTS

3.1. Semi-analytical assessment

In Section 2.2, we computed the probability for true positive evidence for H_1 and $H_{\rm null}$, respectively (Equations 6, 7). Figure 3 shows how these evidences are distributed for sample sizes 10 and 100, and how likely we are to obtain strong evidence $(BF_{H_i,H_j} > 10)$ in the agnostic case where we draw $f_{\rm life}$ from a log-uniform distribution. For n = 10, strong true evidence for H_1 ($H_{\rm null}$) can be expected in $\sim 3\%$ ($\sim 6\%$) of all random experiments. In the majority of cases, the outcome of the survey will be inconclusive. The situation improves with larger samples: for n = 100, 14% (16%) of random samples permit conclusive inference (strong true evidence) under H_1 ($H_{\rm null}$).

The expected resulting evidence further depends on the a priori unknown rate of life's emergence and persistence $f_{\rm life}$ and on the NUV flux threshold. Figure 4 illustrates this dependency: For very low values of either parameter, samples drawn under the null or alternative hypotheses are indistinguishable and the Bayesian evidence is always low. Both higher $f_{\rm life}$ and higher NUV flux thresholds increase the probability of obtaining strong evidence. Larger sample sizes enable this at lower values of these parameters.

So far, we have drawn random values from uni-460 form distributions for $F_{\text{NUV,min}}$, and F_{NUV} , and 461 from a log-uniform distribution for f_{life} . A high 462 biosignature detection rate $f_{\rm life}$ increases the evidence 463 (see Equation 6) but a survey strategy cannot influence 464 it. The same is true for $F_{\text{NUV,min}}$, where again higher 465 values increase the evidence as the binomial distribution 466 for H_1 gets increasingly skewed and shifted away from 467 the one for H_{null} . However, one might select exoplanets 468 for which a biosignature test is performed based on a469 priori available contextual information (Catling et al. 470 2018b) in order to maximize the science yield of in-471 vesting additional resources. For instance, the distri-472 bution of F_{NUV} in the planet sample can be influenced 473 by the survey strategy, and a targeted sampling ap-474 proach could favor extreme values. We model this by 475 distributing F_{NUV} according to different Beta functions 476 and introduce a selectivity parameter $s \in]-1,1[$ such that $F_{\text{NUV}} \sim Beta(1/10^s, 1/10^s)$. Figure 5 shows how 478 the probability of obtaining true strong evidence for H_1 479 scales with selectivity s. For large samples, a high selec-480 tivity $(s \sim 1)$ can increase the probability of obtaining 481 true strong evidence from $\sim 70\%$ for s=0 (random 482 uniform distribution) to > 90%.

3.2. Survey simulations with Bioverse

With HZ occupancy as a requirement for abiogenesis, and barring selection biases beyond stellar brightness, the host star distribution of inhabited planets in a simulated transit survey is skewed toward later spectral types. For a fixed planet sample size, the fraction of inhabited planets is highest for planets orbiting M dwarfs due to the higher NUV fluxes in the HZ of these stars

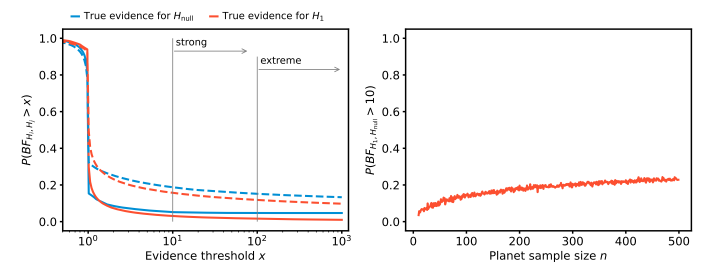


Figure 3. Obtaining true strong evidence with different sample sizes. Left: Probability to reach given evidence levels for H_1 and H_{null} under sample sizes n = 10 (solid) and n = 100 (dashed). Vertical lines denote thresholds for "strong" evidence, $BF_{H_i,H_j} > 10$, and "extreme" evidence, $BF_{H_i,H_j} > 100$. Right: Probability of obtaining true strong evidence for H_1 as a function of sample size n.

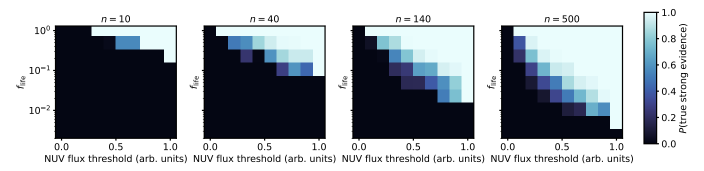


Figure 4. Probability of obtaining true strong evidence for different abiogenesis rates, NUV flux thresholds, and sample sizes. For each of these parameters, higher values increase the probability of yielding strong evidence.

491 (see Figures 2, 6). Their NUV fluxes are generally high492 est at early times ≤ 100 Myr. These host stars, in par493 ticular late subtypes, also provide extended periods of
494 increased NUV emission that overlap with times when
495 some of these planets occupy the HZ (see Figure 2), our
496 requirement for abiogenesis (see Equation 8). Thus −
497 under the UV Threshold Hypothesis − most inhabited
498 transiting planets in the sample orbit M dwarfs.

Here, we are interested in the statistical power of a transit survey with a plausible sample selection and size. In the following, we fix the sample size to 250 and consider two different survey strategies targeting FGK and M dwarfs, respectively. We further investigate the sensitivity of the survey to the a priori unknown threshold NUV flux $F_{\rm NUV,min}$ and the probability of life emerging and persisting $f_{\rm life}$.

3.2.1. Selectivity of simulated transit surveys

In Section 3.1, we showed that the probability of obtaining true strong evidence for the hypothesis that life only originates on planets with a minimum past NUV flux is sensitive to the distribution of sampled past NUV fluxes, i.e., the selectivity of the survey (compare Figure 5). For both surveys targeting M dwarfs and those targeting FGK dwarfs, the maximum NUV distribution first is rather unimodal. Applying the approach from Section 3.1 of fitting a Beta function to the distribution, we find rather low selectivities (see Figure 6), which is likely detrimental for statistical hypothesis tests.

3.2.2. Expected biosignature pattern

A representative recovery of the injected biosignature pattern is shown in Figure 6. There, we assumed an bi22 abiogenesis rate of $f_{\rm life}=0.8$ and a minimum NUV flux for $F_{\rm NUV,min}=300.0\,{\rm erg\,s^{-1}\,cm^{-2}}$. All injected biosignatures are assumed to be detected without false positive ambiguity, and the maximum NUV flux is estimated

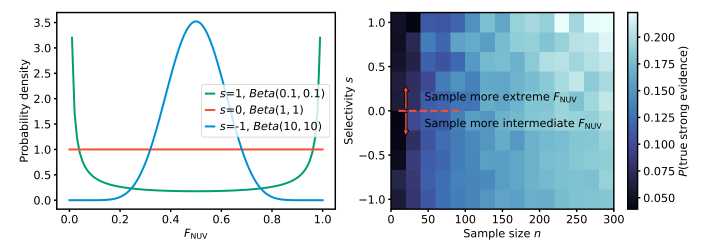


Figure 5. Scaling of the probability of obtaining true strong evidence with sample selectivity. Left: Sampling distribution for different selectivity parameters s. Right: Resulting P(true strong evidence), where f_{life} and $F_{\text{NUV},\text{min}}$ are randomly drawn from log-uniform and uniform distributions, respectively. Sampling more extreme values of F_{NUV} is more likely to yield strong evidence.

567

from the host star's spectral type and age with an uncertainty corresponding to the intrinsic scatter in the NUV fluxes in Richey-Yowell et al. (2023). This leads to a distribution of biosignature detections with detections increasingly occurring above a threshold inferred NUV flux. In this example case, the few biosignature detections in the FGK sample lead to a higher evidence than in the M dwarf sample, where the majority of planets are above the threshold NUV flux.

Figure 7 shows the fraction of inhabited planets under the UV Threshold Hypothesis for different threshold
NUV fluxes and for the limiting case of a probability
for life's emergence and persistence of flife = 1.
This fraction decreases sharply with increasing threshold flux, as fewer planets receive sufficient NUV flux for abiogenesis. Another effect responsible for this drop is that some planets receive the required NUV flux only before entering the HZ – this is especially likely for M dwarfs. For the FGK sample, the fraction of inhabited planets drops at lower threshold fluxes than for the M dwarf sample.

3.2.3. Statistical power and sensitivity to astrophysical parameters

We now investigate the sensitivity of the achieved sta-550 tistical power of our default transit survey to the a pri-551 ori unconstrained threshold NUV flux $F_{\rm NUV,min}$ and the 552 abiogenesis and persistence rate $f_{\rm life}$. Figure 8 shows 553 the statistical power as a function of these parameters 554 for a sample size of N=250. Values of $F_{\rm NUV,min}$ that lie 555 between the extrema of the inferred maximum NUV flux 556 increase the achieved statistical power of the survey, as 557 in this case the dataset under the alternative hypothesis ⁵⁵⁸ H_1 differs more from the null hypothesis. Furthermore, ⁵⁵⁹ higher f_{life} increase the evidence for H_1 .

Parameter space regions with statistical power above 90% lie at $f_{\rm life} > 0.5$ and mostly at threshold NUV fluxes of ~ 200 – $400\,{\rm erg\,s^{-1}\,cm^{-2}}$. Notably, the sensitivity of the M dwarf sample extends into the low NUV flux end due to the broader distribution of maximum past NUV fluxes in this sample. Here, the FGK sample is barely sensitive.

4. DISCUSSION

A key question in the quest to understand the origins of life is which natural processes best explain how living matter spontaneously appears from nonliving matter (e.g., Malaterre et al. 2022). Using astronomical methods, this question will likely not be testable for individual planets but rather for ensembles of planets. The cyanosulfidic scenario for the origins of life (Patel tal. 2015), in particular its predicted existence of a minimum NUV flux required for prebiotic chemistry, offers an opportunity to test an origins of life hypothesis with a statistical transit survey sampling planets with varying NUV flux histories. In the following, we discuss the prospects of testing the UV Threshold Hypothesis in light of our results.

In Section 3.1, we show that testing the UV Threshold Hypothesis suffers from 'nuisance' parameters that hamper inference through astronomical observations. Here, these parameters are the unspecified value of the NUV threshold hypothesized to exist under H_1 , and the unspecified value of a planet $f_{\rm life}$. While the inference of a planet's entire

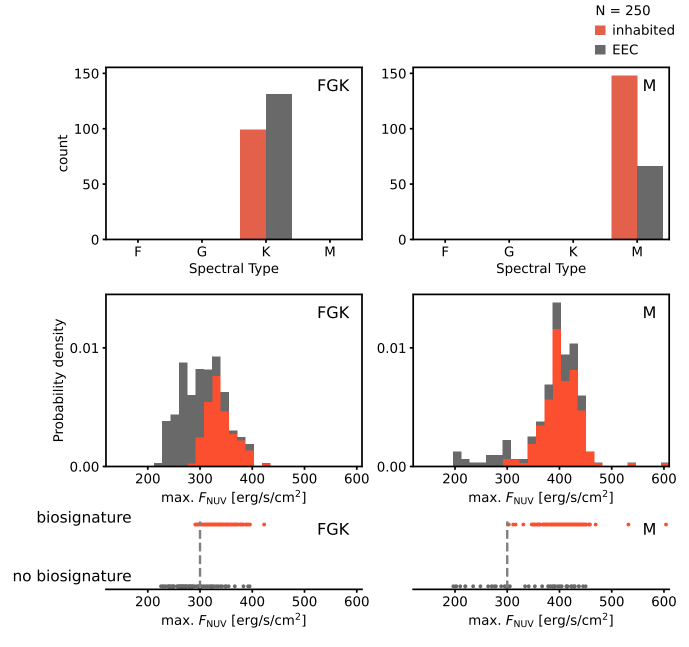


Figure 6. Simulated transit surveys targeting FGK and M stars. Top: Host stars of all transiting EECs and inhabited planets in a simulated transit survey. In the FGK sample, all EECs and all inhabited planets orbit K dwarfs. In an M dwarf sample of the same size, the fraction of inhabited planets is larger. Center: Distribution of inferred maximum past NUV flux in transit surveys targeting EECs around FGK and M stars, respectively. The best-fit beta distributions correspond to selectivities of $s_{\rm FGK} = -0.47$ and $s_{\rm M} = -0.02$. Red areas show inhabited planets for an abiogenesis rate of $f_{\rm life} = 0.8$ and a generic threshold NUV flux $F_{\rm NUV,min} = 300.0\,{\rm erg\,s^{-1}\,cm^{-2}}$. Bottom: Recovered biosignature detections and non-detections of simulated transit surveys. The dashed line denotes $F_{\rm NUV,min}$.

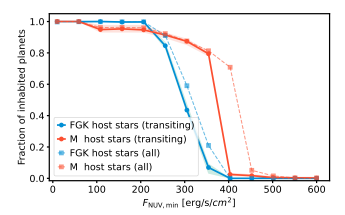


Figure 7. Fraction of inhabited planets for different threshold NUV fluxes under the UV Threshold Hypothesis if the abiogenesis rate $f_{\rm life}=1$. For all samples, the fraction of inhabited planets drops sharply with increasing threshold NUV flux due to the combined effects of never receiving sufficient NUV flux for abiogenesis or receiving it before entering the HZ.

590 UV flux evolution is difficult (e.g., Richey-Yowell et al. 591 2023), the estimated maximum NUV flux that a planet 592 was exposed to may be used as a proxy, at least if one 593 is interested in a minimum threshold flux and makes 594 the assumption that planetary surfaces offer protection 595 against too high UV flux. Indeed, the distribution of the 596 number of planets with detected biosignature in a particular sample of planets with inferred maximum NUV 598 values $F_{\rm NUV}$ depends on both the values of $F_{\rm NUV,min}$ 599 and $f_{\rm life}$ as shown in equation 5.

In our semi-analytical analysis (Section 3.1), we 601 project a possible test performed by a future observer equipped with a sample of exoplanets with derived past maximum NUV exposure for which biosignature detec-604 tion has been attempted. This is necessarily reductive as this observer will have more knowledge about exper-606 imental conditions and will therefore be able to use this 607 information to guide hypothesis testing. For instance, 608 we have made the choice to consider the total number of 609 detected biosignatures as our summary statistic (Equa-610 tion 5), which is not sufficient to infer $F_{NUV,min}$ and ₆₁₁ f_{life} separately. However, by conditioning the Bayes factor to these variables (Equation 6), we calculate the 613 probability distribution of the Bayesian evidence in faof H_1 . In doing so, we may evaluate how evidence 615 depends on the uncertainty over these unknown parame-616 ters in general terms, without assuming which particular 617 test a future observer might actually choose to perform 618 over real data when available. From this, we can see that target selection can strongly affect the conclusiveness of 620 a future test of the UV Threshold Hypothesis.

The particular finding that prioritizing extreme values 622 of past NUV flux can enhance statistical power likely 623 clashes with observational constraints, as the composi-624 tion of the subset of planets that we can observe and 625 for which detection of biosignature can be attempted is 626 not independent from their NUV flux history. Hence, 627 for our future observer, selectivity and sample size may be in conflict. This trade-off can be quantified in terms of expected evidence yield, which we have done in Sec-630 tion 3.1. Our analysis shows that regardless of selec-631 tivity, sample sizes smaller than 50 likely result in in-632 conclusive tests, and that increasing selectivity towards extreme F_{NUV} offers limited inference gains compared 634 to the uniform case (s=0; Figure 5). For larger sam-635 ples, however, a narrow distribution of F_{NUV} may pre-636 vent inference entirely. We thus argue that selecting a 637 sample with F_{NUV} distributed uniformly or emphasiz-638 ing extreme values should – barring any practical coun-639 terarguments – be considered in any future attempt at 640 testing the UV Threshold Hypothesis. Since the prac-641 tical implementation of an exoplanet survey can stand 642 in the way of such a selection, the following discussion focuses on the results of our transit survey simulations with Bioverse.

4.2. How planetary context may constrain the UV Threshold Hypothesis

It comes to no surprise that the success rate for testing 648 the UV Threshold Hypothesis is sensitive to the sample size of the survey and to the occurrence of life on temper-650 ate exoplanets. As we have shown, the statistical power of this test also depends on the distribution of past NUV 652 fluxes in the sample and on the threshold flux. Opti-653 mizing the survey to sample a wide range of NUV flux 654 values, particularly at the extremes, can enhance the 655 likelihood of obtaining strong evidence for or against 656 the hypothesis. Intermediate values of the threshold 657 NUV flux are more likely to yield strong evidence than 658 extreme values, as the dataset under the alternative hypothesis H_1 differs more from the null hypothesis in this case while still being sufficiently populated. The thresh-661 old flux is, of course, a priori unknown and we cannot 662 influence it. If, however, better theoretical predictions 663 for the required NUV flux for abiogenesis become available (Rimmer et al. 2021a), the survey strategy can be further optimized, for instance by targeting planets that are estimated to have received a NUV flux slightly be-667 low and above this threshold or by applying a bisection algorithm in a sequential survey (Fields et al. 2023).

4.3. An M dwarf opportunity

An interesting aspect lies in the distribution of host tar properties, as different spectral types probe differ-

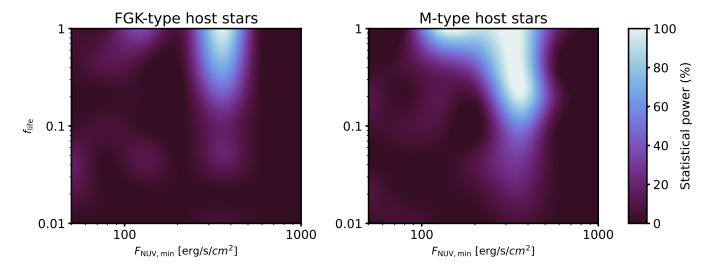


Figure 8. Statistical power as a function of threshold NUV flux and abiogenesis rate. Even for a large sample (here: N=250), a high statistical power of the transit survey requires high **rates of life emerging and persisting** f_{life} . Intermediate values of $F_{\text{NUV,min}}$ are more likely to yield strong evidence than extreme values. For $f_{\text{life}} \gtrsim 80\%$, the sensitivity of the M dwarf sample extends into the low NUV flux end.

ent past NUV flux regimes. FGK stars exhibit a rel-673 atively narrow range of maximum past NUV fluxes 674 in the HZ, which may – depending on the (unknown) 675 threshold flux – limit the diagnostic power of a survey. 676 A pure FGK sample would only be sensitive to flux thresholds in the range of $\sim 200-400\,\mathrm{erg\,s^{-1}\,cm^{-2}}$. A detection of biosignatures in such a sample 679 would likely only suggest that either low NUV 680 fluxes are sufficient for abiogenesis or that an alternative abiogenic pathway may be at play (e.g., Westall et al. 2018). Conversely, a lack of biosignatures in a sufficiently large FGK sample 684 could indicate that the actual threshold is higher than the maximum past flux levels reached by 686 FGK hosts. We further note that under the 687 UV Threshold Hypothesis, the fraction of inhab-688 ited planets in an FGK sample declines rapidly 689 for increasing threshold fluxes, as shown in Fig-690 ure 7. In addition, if the timescale for the emergence of life is $\gg 1 \,\mathrm{Myr}$, the fraction of inhabited 692 planets in an FGK sample may be negligible (see 693 Section C).

On the other hand, M dwarfs show a wider distribution of maximum past NUV fluxes in their HZs. While old M dwarfs can be considered low-UV environments, a significant fraction of them emit high NUV fluxes into their HZ during their early stages, in particular later subtypes (Richey-Yowell et al. 2023). This will help to to test the high NUV flux end of the UV Threshold Hypothesis; a higher occurrence of biosignatures here would support the hypothesis that a higher NUV flux is favorable or necessary for life. At the same time, a fraction of host stars in our M dwarf sample extends it to
lower maximum past NUV fluxes, enabling tests of the
low NUV flux end of the hypothesis. The higher and
more variable NUV fluxes in M dwarfs thus increase the
likelihood of obtaining strong evidence for or against the
UV Threshold Hypothesis.

The combination of a lack of UV radiation today, which makes biosignature gases more detectable (Segura et al. 2005), and a UV-rich past that may have enabled abiogenesis could make M dwarfs the preferred targets for biosignature searches. We note that relevant mission concepts, such as the Large Interferometer for Exoplanets (LIFE, Quanz et al. 2022; Glauser et al. 2024), include M-dwarf systems among their primary targets (Kammerer & Quanz 2018; Carrión-González et al. 2023). Our findings underscore the importance of constraining the UV emission profiles of EEC host stars throughout their evolutionary stages to assess the viability of M-dwarf planets as testbeds for theories on the origins of life (Rimmer et al. 2021a; Ranjan et al. 2023).

4.4. Sensitivity to astrophysical parameters

Our Bioverse simulations that take into account expolared demographics, the evolution of habitability and NUV fluxes, and observational biases show that not only the likelihood of a conclusive test of the UV Threshold Hypothesis, but also the likelihood of successful biosignature detection itself is extremely sensitive to the threshold NUV flux if the hypothesis is true. Even if all biosignatures can be detected and the nominal rate of life's emergence and persistence is very high, say $f_{\rm life} = 1$, under the condition that prebiotic chemistry

753

782

736 requires a minimum NUV flux and liquid water, if the 737 threshold flux turns out to be high the probability of 738 finding life on a randomly selected planet may be very 739 low. As we showed, for high required fluxes the two re-740 quirements of simultaneous HZ occupancy and sufficient 741 NUV flux conspire to diminish the fraction of inhabited 742 planets in the sample. Taking the inferred fluxes from Richey-Yowell et al. (2023) at face value (but taking into account intrinsic scatter), a minimum required NUV flux 745 of $\gtrsim 400\,\mathrm{erg\,s^{-1}\,cm^{-2}}$ reduces the fraction of inhabited 746 planets to below $\sim 1\%$. This not only calls for a large 747 sample size and a targeted sample selection preferring 748 high expected past NUV fluxes, but also highlight the 749 necessity of continued theoretical and experimental re-750 search into the role of UV radiation in prebiotic chem-751 istry (Ranjan et al. 2017b; Rimmer et al. 2018, 2021a).

4.5. Contextual support for potential biosignature detections

The predicted interplay of NUV flux and HZ occurospancy in enabling abiogenesis via the cyanosulfidic scenario could in principle be used to add or remove credibility from a tentative biosignature detection. For example, with a strong prior belief that this scenario is the only viable one for the origins of life (see Section 4.7.6), a biosignature detection on a planet orbiting a strongly UV-radiating star may add credibility to the detection. Conversely, a biosignature detection on a planet estimated to have received very little UV radiation would increase the likelihood of a false positive detection. On the other hand, should the detection in the latter case be confirmed, it could be used to falsify the UV Threshold Hypothesis.

Our simulations find no clear criterion for the credibility of a biosignature detection based on spectral type of the host star, as both FGK and M dwarf samples show similar maximum past NUV flux distributions. While there is a significant preference for M dwarfs in the case of a long timescale for abiogenesis (see Appendix C), the fraction of inhabited planets in the M dwarf sample drops at similar threshold fluxes as in the FGK sample if the timescale for abiogenesis is short (see Section 3.2). A potentially inhabited planet's host star spectral type may thus not be a strong indicator for the credibility of a biosignature detection in the context of the UV Threshold Hypothesis.

4.6. Overall prospects for testing the UV Threshold Hypothesis

Our results show that the UV Threshold Hypothesis is testable with potential future exoplanet surveys, but

that the success of such a test depends on the sample size, the distribution of past NUV fluxes, and several rms unknown astrophysical nuisance parameters. Even ungerial deridealized conditions, obtaining strong evidence for or against the hypothesis likely requires sample sizes on the order of 100 (see Section 3.1). This is true for a future transit survey, the specifics of which we have reflected in our Bioverse simulations (see Section 3.2). However, we have shown that the impacts from the combined requirements of the UV Threshold Hypothesis on the fraction of inhabited planets in a sample are comparable in the non-transiting case.

Given the challenging nature of detecting and charac-799 terizing small (Earth-sized) exoplanets, most exoplanet mission concepts currently under development or consid-801 ered lack the potential for characterizing large enough 802 samples. Ground-based 25–40-meter class extremely 803 large telescopes are expected to have the capabilities 804 to detect biosignatures on exoplanets like Proxima Censos tauri b (e.g., Wang et al. 2017; Hawker & Parry 2019; 806 Zhang et al. 2024; Vaughan et al. 2024). Hardegree-807 Ullman et al. (2025) used Bioverse to determine posos tential yields for a 10-year direct imaging and highresolution spectroscopy survey of O₂ on the Giant Mag-810 ellan Telescope (GMT) and on the Extremely Large ** Telescope (ELT) and found that between 7 and 19 habit-812 able zone Earth-sized planets could be probed for Earth-813 like oxygen levels. Such a sample is too small to test 814 the UV Threshold Hypothesis, but it may be synergis-815 tic with other detection methods.

The Habitable Worlds Observatory (HWO) is expected to characterize a sample of ~ 25 Earth analogs (Mamajek & Stapelfeldt 2023; Tuchow et al. 2024). Depending on the technical design, LIFE is expected to target 25–80 EECs (Kammerer & Quanz 2018; Quanz et al. 2022), which could be just sufficient to constrain the UV Threshold Hypothesis.

If the rate of life's emergence and persistence $f_{\rm life}$ is at a 1% level or lower, no currently ensus visioned future exoplanet mission has projected sample sizes sufficient to test the UV Thresheld Hypothesis. One possible exception is the Nausilus Space Observatory concept (Apai et al. 2019, 2022). Nautilus aims to characterize up to ~ 1000 EEC via transmission spectroscopy, building on an innovative opaitical technology. To guide the definition of future biosignature surveys, it is important to refine predictions on the role of UV radiation in prebiotic chemistry with both theoretical and experimental work.

Our work is based on a number of assumptions and simplifications that may affect the results and conclusions. We discuss some of these caveats here.

839

4.7.1. The UV Threshold Hypothesis as a narrow step function

A key aspect of the UV Threshold Hypothesis is is the proposed step-function dependence of abiogenesis likelihood on UV flux. This approach stems from the constraints governing photo-chemical pathways, which exhibit a threshold behavior: below a certain flux, competing thermal reactions dominate, preventing abiogenesis, while above the threshold, UV photochemistry proceeds at sufficient rates, and other stochastic processes become rate-limiting.

Ranjan et al. (2017b) speculated that UV pho-852 853 tochemistry might be rate-limiting for abiogen-854 esis, particularly on planets orbiting M dwarfs, 855 due to their lower baseline UV fluxes. This could 856 delay abiogenesis by orders of magnitude, result-857 ing in a continuous dependence of abiogenesis 858 likelihood on UV flux. However, recent stud-859 ies challenge this view for the cyanosulfidic sce-860 nario. Rimmer et al. (2021a) calculated photo-861 chemical timescales on early Earth at 180-300 862 hours (7.5–12.5 days), significantly shorter than 863 the timescale for stochastic geological events also required for the scenario (Rimmer 2023). 865 Even with 1000x slower photochemistry on M-866 dwarf planets, prebiotic photochemistry like sul-867 fite photolysis would still occur on a geologically ** negligible timescale of 20-30 years, meaning that s69 the photochemistry is unlikely to be rate limit-870 ing compared to stochastic geological processes, 371 justifying the step-function model.

Nonetheless, alternative abiogenesis pathways or combinations of pathways may exhibit contin10 uous or mixed dependencies on UV flux. While the step-function formalism is justified for the cyanosulfidic scenario, future work should explore UV dependencies across other scenarios to refine predictions for biosignature distributions and testable hypotheses.

4.7.2. Existence of an atmosphere-crust interface

By its nature, cyanosulfidic scenario relies on rock surfaces exposed to the planetary atmosphere. Water worlds that have their entire planetary surface covered by oceans contradict this requirement and do not also low for the wet-dry cycling inherent to this origin of life see scenario. The competition of tectonic stress with grav-

s87 itational crustal spreading (Melosh 2011) sets the max888 imum possible height of mountains, which in the solar
889 system does not exceed ~20 km. Such mountains will
890 be permanently underwater on water worlds. Another
891 impediment to wet-dry cycles may be tidal locking of
892 the planet as it stalls stellar tide-induced water move893 ment and diurnal irradiation variability (e.g., Ranjan
894 et al. 2017b). However, recent dynamical models sug895 gest tidally locked planets to undergo rapid drift of their
896 sub-stellar point (Revol et al. 2024).

4.7.3. Stellar flares

Our assumptions on past UV flux neglect the contrisue bution of stellar flares, which may be hypothesized as an alternative source of UV light (Buccino et al. 2007; Rangon et al. 2017a). This concerns mainly ultracool dwarfs, due to their low quiescent emission and high pre-main sequence stellar activity (Buccino et al. 2007; West et al. 2008). However, recent work indicates that the majority of stars show inadequate activity levels for a sufficient contribution through flares (Glazier et al. 2020; Ducrot et al. 2020; Günther et al. 2020).

4.7.4. Atmosphere transmission

We do not take into account absorption of UV radia-910 tion by the planetary atmosphere. Theoretical work suggests that the atmosphere of prebiotic Earth was largely 912 transparent at NUV wavelengths with the only known 913 source of attenuation being Rayleigh scattering (Ran-914 jan & Sasselov 2017; Ranjan et al. 2017a). We thus 915 approximated surface UV flux using top-of-atmosphere 916 fluxes. If there are planets in a sample that do not have 917 a transparent atmosphere at NUV wavelengths and re-918 quire higher fluxes for abiogenesis, the fraction of inhab-919 ited planets in the sample will be lower. However, these 920 planets will not pollute the below-threshold subsample, 921 as they will not be able to host life under the UV Thresh-922 old Hypothesis. Exoplanet surveys focusing on highly 923 irradiated planets offer an opportunity to constrain the 924 typical oxidation state of rocky exoplanets, providing 925 insights into the average composition of their secondary 926 atmospheres (Lichtenberg & Miguel 2024). This is par-927 ticularly relevant for prebiotic worlds, as varying oxi-928 dation states significantly perturb the classical habit-929 able zone concept (Nicholls et al. 2024) and also influ-930 ence surface UV levels through changing atmospheric 931 transmission. Optimally, the atmospheric composition 932 of young rocky protoplanets will be probed to constrain 933 the possible range of atmospheric and mantle oxidation 934 states during early planetary evolution by future direct 935 imaging concepts (Cesario et al. 2024).

937 4.7.5. Other mechanisms regulating habitability and abiogenesis

Our study focuses primarily on two factors that 940 may regulate the emergence and persistence of 941 life on a planet: the NUV flux and the via-942 bility of liquid water, which provide a testable 943 framework for assessing biosignature distribu-944 tions. Of course, planetary habitability and abio-945 genesis depend on a broader range of physical 946 and chemical conditions. For one, a planet must 947 retain an atmosphere capable of supporting liq-948 uid water and surface chemistry. Atmospheric 949 escape processes have been studied in detail, and 950 their occurrence is supported by planet forma-951 tion models (e.g., Owen & Wu 2013; Schlicht-952 ing et al. 2014; Ginzburg et al. 2016; Mordasini 953 2020; Burn et al. 2024) as well as exoplanet de-954 mographics (e.g., Owen & Estrada 2019; Berg-955 sten et al. 2022; Rogers et al. 2021). Popula-956 tion synthesis studies demonstrate that models 957 of atmospheric escape can explain key statisti-958 cal features of the observed exoplanet popula-959 tion (Rogers & Owen 2020; Emsenhuber et al. 960 2021; Schlecker et al. 2021; Burn et al. 2024).

Planets around M dwarfs, in particular, may 962 be subject to significant atmospheric loss due to 963 the extended high-luminosity pre-main-sequence 964 phase of their host stars (e.g., Luger & Barnes 965 2015). Recent works (e.g., Coy et al. 2024; Luque 966 et al. 2024) interpret JWST eclipse measurements as growing evidence for the absence of sub-968 stantial atmospheres on M-dwarf rocky exoplanets. If these planets indeed lack atmospheres, 970 their habitability would be primarily dictated by 971 the rate of atmospheric escape (e.g., Owen & 972 Campos Estrada 2020). However, alternative interpretations remain viable: Ducrot et al. (2024) 974 and Hammond et al. (2025) recently demon-975 strated that current JWST data cannot defini-976 tively distinguish between a bare-rock scenario ₉₇₇ and an atmosphere composed primarily of N_2 -978 CO₂-H₂O. This underscores the need for future 979 observations and modeling efforts to constrain ⁹⁸⁰ atmospheric retention and escape processes more 981 robustly.

Beyond atmospheric escape, internal heating from tidal forces or radioactive decay can extend or constrain the limits of planetary surface habitability (e.g., Barnes et al. 2013; Oosterloo et al. 2021). Tidal effects are especially relevant for HZ M dwarf planets, where strong stellar intersactions can drive internal heating, loss of water

or an atmosphere, or runaway greenhouse conditions. Tidal locking may also create extreme climate zones, challenging habitability. While not modeled here, the factors outlined above may offer directions for future work.

4.7.6. Bayesian Evidence and the Influence of Priors

Our semi-analytical analysis (Section 3.1) employs Bayes factors to quantify the evidence in favor of or against the UV Threshold Hypothesis that future observations may provide.

Using the Bayes theorem to estimate posterior probabilities would allow for a more complete assessment by quantifying information gain
and integrating prior knowledge into hypothesis testing. In practice, implementation of a full
Bayesian inference is often hindered by the subestivity of prior distributions. Here, we thus
chose to provide merely a general assessment of
the potential evidence yield from future observations. In essence, we addressed the question: By
how much does a particular observation tip the
scale between the UV Threshold Hypothesis and
the null hypothesis?

5. CONCLUSIONS

We propose that specific origins-of-life scenarios may loss leave a detectable imprint on the distribution of biosignatures in exoplanet populations. We have investigated the potential of upcoming exoplanet surveys to test the hypothesis — motivated by the cyanosulfidic origins-of-lose life scenario — that a minimum past NUV flux is required for abiogenesis. To this end, we first employed a semi-lose analytical Bayesian analysis to estimate probabilities of obtaining strong evidence for or against this hypothe-lose sis. We then used the Bioverse framework to assess lose the diagnostic power of realistic transit surveys, taking into account exoplanet demographics, time-dependency of habitability and NUV fluxes, observational biases, and target selection.

Our main findings are:

1013

1028

1029

1030

1031

1033

1034

1035

1036

1037

- 1. The UV Threshold Hypothesis of the cyanosulfidic scenario for the origins of life should lead to a correlation between past NUV flux and current occurrence of biosignatures that may be observationally testable.
- 2. The required sample size for detecting this correlation depends on the abiogenesis rate on temperate exoplanets and the distribution of host star properties in the sample; in particular their maximum

past NUV fluxes. Samples smaller than 50 planets are unlikely to yield conclusive results.

1073

1074

1120

1038

1039

1040

1041

1042

1043

1044

1045

1047

1048

1049

1050

1051

- 3. Under the UV Threshold Hypothesis, the fraction of inhabited planets in a transit survey is sensitive to the threshold NUV flux and is expected to drop sharply for required fluxes above a few hundred ${\rm erg \, s^{-1} \, cm^{-2}}$.
- 4. If the predicted UV correlation exists, obtaining strong evidence for the hypothesis is likely (≥ 80%) for sample sizes ≥ 100 if the abiogenesis rate is high (≥ 50%) and if no very high NUV fluxes are required. A survey strategy that targets extreme values of inferred past NUV irradiation increases the diagnostic power.
- 5. Samples of planets orbiting M dwarfs overall yield higher chances of successfully testing the UV Threshold Hypothesis. They may also be more likely to yield biosignature detections under this hypothesis, in particular if the origins timescale is long.

Overall, our work demonstrates that future exoplanet surveys have the potential to test the hypothesis that most a minimum past NUV flux is required for abiogenesis. More generally, we found that models of the origins of life provide hypotheses that may be testable with these surveys. Conducting realistic survey simulations with representative samples is important to identify testable science questions, support trade studies, help define science cases for future missions, and guide further theorem oretical and experimental work on the origins of life. Our work highlights the importance of understanding the context in which a biosignature detection is made, which can not only help to assess the credibility of the detection but also to test competing hypotheses on the origins of life on Earth and beyond.

ACKNOWLEDGMENTS

The authors thank Kevin Heng, Dominik Hintz, Do-1075 1076 minika Itrich, Chia-Lung Lin, and Rhys Seeburger for 1077 insightful discussions. We thank the anonymous referee 1078 for providing constructive critical feedback that helped 1079 to improve this manuscript. This material is based upon 1080 work supported by the National Aeronautics and Space 1081 Administration under Agreement No. 80NSSC21K0593 1082 for the program "Alien Earths". The results reported 1083 herein benefited from collaborations and/or informa-1084 tion exchange within NASA's Nexus for Exoplanet 1085 System Science (NExSS) research coordination net-1086 work sponsored by NASA's Science Mission Direc-1087 torate. This work has made use of data from the Euro-1088 pean Space Agency (ESA) mission Gaia (https://www. 1089 cosmos.esa.int/gaia), processed by the Gaia Data Pro-1090 cessing and Analysis Consortium (DPAC, https://www. 1091 cosmos.esa.int/web/gaia/dpac/consortium). Funding 1092 for the DPAC has been provided by national institutions, in particular the institutions participating in the 1094 Gaia Multilateral Agreement. T.L. was supported by the Branco Weiss Foundation, the Netherlands eScience 1096 Center (PROTEUS project, NLESC.OEC.2023.017), and the Alfred P. Sloan Foundation (AEThER project, G202114194).

AUTHOR CONTRIBUTIONS

M.S., D.A., and S.R. conceived the project, planned 1101 its implementation, and interpreted the results. M.S. 1102 developed the planetary evolution component to 1103 Bioverse, carried out the hypothesis tests and statis-1104 tical analyses, and wrote the manuscript. D.A. leads 1105 the "Alien Earths" program through which this project 1106 is funded, helped to guide the strategy of the project, and provided text contributions. A.A. carried out the 1108 semi-analytical computations regarding the correlation 1109 of past UV flux and biosignature occurrence. S.R. advised on planetary NUV flux evolution and the cyanosul-1111 fidic scenario of the origins of life. R.F. wrote the initial 1112 draft of the Introduction and advised on the evolutionary biology aspects of the project. K.H.-U. contributed 1114 to the Bioverse software development and simulations. 1115 T.L. supported the selection of testable hypotheses and provided text contributions to the initial draft. S.M. advised on the scope of the project and supported the 1118 selection of testable hypotheses. All authors provided 1119 comments and suggestions on the manuscript.

REPRODUCIBILITY

1125

1126

1121 All code required to reproduce our results, figures, 1122 and this article itself is available at https://github.com/1123 matiscke/originsoflife.

APPENDIX

A. PHOTON NUMBER FLUXES AND ENERGY FLUXES

The NUV fluxes we used in our simulations are 1127 given in units of energy flux, i.e., erg s⁻¹ cm⁻², 1129 due to their availability across different spectral 1130 types and evolutionary stages of stars. Here, we assess the robustness of our conclusions when 1132 considering photon number fluxes instead. 1133 first computed blackbody spectral energy distri-1134 butions (SEDs) for representative effective tem-1135 peratures corresponding to K-type, early M-1136 type, and late M-type stars, and across the evo-1137 lutionary stages considered in the main text. 1138 While blackbody SEDs are not necessarily repre-1139 sentative of M-dwarf SEDs in the NUV (Seager 1140 et al. 2013; Rugheimer et al. 2015), the spec-1141 trally resolved evolution of M-dwarf SEDs as 1142 a function of age is not known; we therefore 1143 adopt this simplified prescription for the pur-1144 pose of this sensitivity test. We then normalized 1145 these SEDs to align with the observed UV fluxes 1146 from Richey-Yowell et al. (2023) and used them

to calculate mean number flux densities (phonom tons $\rm cm^{-2}\,s^{-1}\,nm^{-1})$ in the 200–280 nm wavelength range.

We found that the derived flux densities for all considered spectral types exceed the threshold estimated by Rimmer et al. (2018) during earlier stages of stellar evolution (Figure 9). Notably as in the case of energy fluxes – later spectral types exhibit higher maximum past NUV num1156 ber flux densities.

We then calculated the ratio of the maximum past NUV fluxes in K dwarfs and late M dwarfs in both photon number flux and energy flux regimes. We found that this "dynamic range" is slightly larger in the photon number flux regime (2.55) compared to the energy flux regime (2.45). This suggests that our use of energy flux may slightly underestimate the variations in NUV exposure across different stellar types, rendering our conclusions regarding the impact of host star spectral type on the UV Threshold Hypothesis conservative. Our results are thus robust when transitioning between energy flux and photon number flux representations.

REFERENCES

```
1171 Affholder, A., Mazevet, S., B., S., Apai, D., & Ferrière, R.
      2025, The Astronomical Journal,
1172
      doi: 10.3847/1538-3881/ada384
1174 Alfeld, P. 1984, Computer Aided Geometric Design, 1, 169,
      doi: 10.1016/0167-8396(84)90029-3
1175
1176 Apai, D., Milster, T. D., Kim, D. W., et al. 2019, The
      Astronomical Journal, 158, 83,
1177
      doi: 10.3847/1538-3881/ab2631
1178
1179 Apai, D., Milster, T. D., Kim, D., et al. 2022, in Optical
      Manufacturing and Testing XIV, Vol. 12221 (SPIE),
1180
      59-71, doi: 10.1117/12.2633184
1181
Baraffe, I., Chabrier, G., Allard, F., & Hauschildt, P. H.
      1998, Astronomy and Astrophysics, v.337, p.403-412
1183
      (1998), 337, 403
1184
1185 Barnes, R., Mullins, K., Goldblatt, C., et al. 2013,
      Astrobiology, 13, 225, doi: 10.1089/ast.2012.0851
1186
```

```
1187 Baross, J. A., & Hoffman, S. E. 1985, Origins of life and
      evolution of the biosphere, 15, 327,
      doi: 10.1007/BF01808177
1190 Bergsten, G. J., Pascucci, I., Mulders, G. D., Fernandes,
      R. B., & Koskinen, T. T. 2022, The Astronomical
      Journal, 164, 190, doi: 10.3847/1538-3881/ac8fea
1192
Bixel, A., & Apai, D. 2021, The Astronomical Journal, 161,
      228, doi: 10.3847/1538-3881/abe042
1195 Bonati, I., & Ramirez, R. M. 2021, Monthly Notices of the
      Royal Astronomical Society, 504, 1029,
      doi: 10.1093/mnras/stab891
   Brasier, M. D., Matthewman, R., McMahon, S., & Wacey,
      D. 2011, Astrobiology, 11, 725, doi: 10.1089/ast.2010.0546
Brunner, E., & Munzel, U. 2000, Biometrical Journal, 42,
      17, doi: 10.1002/(SICI)1521-4036(200001)42:1<17::
      AID-BIMJ17>3.0.CO;2-U
1203 Buccino, A. P., Lemarchand, G. A., & Mauas, P. J. D.
      2007, Icarus, 192, 582, doi: 10.1016/j.icarus.2007.08.012
```

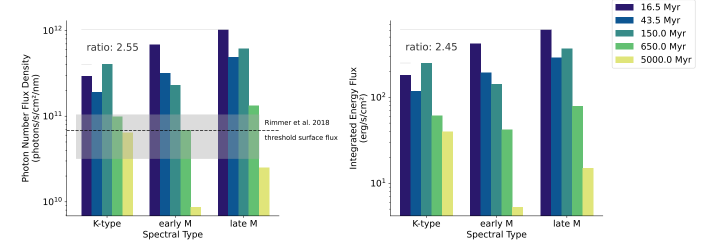


Figure 9. Comparison of NUV flux evolution in terms of photon number flux and energy flux for K-type, early M-type, and late M-type stars. In both regimes, later spectral types exhibit higher maximum values. The ratio of the maximum past NUV fluxes in K dwarfs and late M dwarfs is slightly larger in the photon number flux regime compared to the energy flux regime, suggesting that our conclusions regarding the impact of host star spectral type on the UV Threshold Hypothesis are conservative.

```
1236 Emsenhuber, A., Mordasini, C., Burn, R., et al. 2021,
1205 Burn, R., Mordasini, C., Mishra, L., et al. 2024, Nature
      Astronomy, 8, 463, doi: 10.1038/s41550-023-02183-7
                                                                          Astronomy & Astrophysics, 656, A70,
1206
                                                                    1237
    Carrión-González, Ó., Kammerer, J., Angerhausen, D.,
                                                                          doi: 10.1051/0004-6361/202038863
1207
                                                                    1238
      et al. 2023, Astronomy & Astrophysics, 678, A96,
                                                                        Fields, B., Gupta, S., & Sandora, M. 2023, International
1208
                                                                    1239
      doi: 10.1051/0004-6361/202347027
                                                                          Journal of Astrobiology, 22, 583,
                                                                    1240
1209
    Catling, D. C., Krissansen-Totton, J., Kiang, N. Y., et al.
1210
                                                                          doi: 10.1017/S1473550423000150
      2018a, Astrobiology, 18, 709, doi: 10.1089/ast.2017.1737
                                                                        Fox, S., & Strasdeit, H. 2013, Astrobiology, 13, 578,
1211
    Catling, D. C., Krissansen-Totton, J., Kiang, N. Y., et al.
1212
                                                                          doi: 10.1089/ast.2012.0934
                                                                    1243
      2018b, Astrobiology, 18, 709, doi: 10.1089/ast.2017.1737
                                                                        Ginzburg, S., Schlichting, H. E., & Sari, R. 2016, The
                                                                    1244
    Cesario, L., Lichtenberg, T., Alei, E., et al. 2024, Large
                                                                           Astrophysical Journal, 825, 29,
                                                                    1245
      Interferometer For Exoplanets (LIFE). XIV. Finding
1215
                                                                          doi: 10.3847/0004-637x/825/1/29
                                                                    1246
      Terrestrial Protoplanets in the Galactic Neighborhood,
1216
                                                                        Glauser, A. M., Quanz, S. P., Hansen, J., et al. 2024, in
                                                                    1247
      arXiv, doi: 10.48550/arXiv.2410.13457
1217
                                                                          Optical and Infrared Interferometry and Imaging IX, Vol.
                                                                    1248
      Chaverot, G., Turbet, M., Bolmont, E., & Leconte, J. 2022,
1218
                                                                           13095 (SPIE), 354-374, doi: 10.1117/12.3019090
                                                                    1249
      Astronomy & Astrophysics, 658, A40,
1219
                                                                        Glazier, A. L., Howard, W. S., Corbett, H., et al. 2020, The
                                                                    1250
      doi: 10.1051/0004-6361/202142286
1220
                                                                           Astrophysical Journal, 900, 27,
                                                                    1251
      loy, B. P., Ih, J., Kite, E. S., et al. 2024, Population-Level
1221
                                                                          doi: 10.3847/1538-4357/aba4a6
                                                                    1252
      Hypothesis Testing with Rocky Planet Emission Data: A
1222
                                                                        Günther, M. N., Zhan, Z., Seager, S., et al. 2020, The
                                                                    1253
      Tentative Trend in the Brightness Temperatures of
1223
                                                                           Astronomical Journal, 159, 60,
                                                                    1254
      M-Earths, arXiv, doi: 10.48550/arXiv.2412.06573
                                                                          \mathbf{doi:}\ 10.3847/1538\text{--}3881/ab5d3a
                                                                    1255
    Deamer, D., Damer, B., & Kompanichenko, V. 2019,
                                                                         Hammond, M., Guimond, C. M., Lichtenberg, T., et al.
                                                                    1256
      Astrobiology, 19, 1523, doi: 10.1089/ast.2018.1979
                                                                           2025, The Astrophysical Journal Letters, 978, L40,
                                                                    1257
1227 Deamer, D. W., & Georgiou, C. D. 2015, Astrobiology, 15,
                                                                          doi: 10.3847/2041-8213/ada0bc
                                                                    1258
      1091, doi: 10.1089/ast.2015.1338
                                                                        Hardegree-Ullman, K. K., Apai, D., Bergsten, G. J.,
Dodd, M. S., Papineau, D., Grenne, T., et al. 2017, Nature,
                                                                    1259
                                                                          Pascucci, I., & López-Morales, M. 2023, The
      543, 60, doi: 10.1038/nature21377
                                                                    1260
1230
                                                                          Astronomical Journal, 165, 267,
Ducrot, E., Gillon, M., Delrez, L., et al. 2020, Astronomy &
                                                                    1261
                                                                          doi: 10.3847/1538-3881/acd1ec
      Astrophysics, 640, A112,
                                                                    1262
1232
                                                                    1263 Hardegree-Ullman, K. K., Apai, D., Haffert, S. Y., et al.
      doi: 10.1051/0004-6361/201937392
1233
                                                                          2025, The Astronomical Journal, 169, 171,
    Ducrot, E., Lagage, P.-O., Min, M., et al. 2024, Nature
1234
                                                                    1264
       Astronomy, 1, doi: 10.1038/s41550-024-02428-z
                                                                          doi: 10.3847/1538-3881/adb02f
                                                                    1265
```

```
1266 Hawker, G. A., & Parry, I. R. 2019, MNRAS, 484, 4855,
                                                                   1316 Mulkidjanian, A. Y., Bychkov, A. Y., Dibrova, D. V.,
                                                                         Galperin, M. Y., & Koonin, E. V. 2012, Proceedings of
      doi: 10.1093/mnras/stz323
1267
                                                                   1317
1268 Ingersoll, A. P. 1969, Journal of Atmospheric Sciences, 26,
                                                                         the National Academy of Sciences, 109, E821,
                                                                   1318
      1191, doi: 10.1175/1520-0469(1969)026<1191:
                                                                         doi: 10.1073/pnas.1117774109
                                                                   1319
1269
      TRGAHO>2.0.CO;2
                                                                       Nicholls, H., Lichtenberg, T., Bower, D. J., &
                                                                   1320
1270
    Jeffreys, H. 1939, Theory of Probability
                                                                         Pierrehumbert, R. 2024, Magma Ocean Evolution at
                                                                   1321
1271
1272 Kammerer, J., & Quanz, S. P. 2018, Astronomy &
                                                                         Arbitrary Redox State, arXiv,
                                                                   1322
      Astrophysics, 609, A4, doi: 10.1051/0004-6361/201731254
                                                                         doi: 10.48550/arXiv.2411.19137
                                                                   1323
1273
1274 Kasting, J. F. 1991, Icarus, 94, 1,
                                                                   Nielson, G. M. 1983, Mathematics of Computation, 40, 253,
      doi: 10.1016/0019-1035(91)90137-I
                                                                         doi: 10.1090/S0025-5718-1983-0679444-7
    Kasting, J. F., Whitmire, D. P., & Reynolds, R. T. 1993,
                                                                   1326 Noack, L., Höning, D., Rivoldini, A., et al. 2016, Icarus,
1276
      Icarus, 101, 108, doi: 10.1006/icar.1993.1010
                                                                         277, 215, doi: 10.1016/j.icarus.2016.05.009
1278 Kipping, D. 2021, Proceedings of the National Academy of
                                                                   1328 Oosterloo, M., Höning, D., Kamp, I. E. E., & van der Tak,
      Sciences, 118, e2026808118, doi: 10.1073/pnas.2026808118
                                                                         F. F. S. 2021, doi: 10.1051/0004-6361/202039664
   Kite, E. S., & Ford, E. B. 2018, The Astrophysical Journal,
                                                                   1330 Owen, J., & Estrada, B. C. 2019, 12, 1
      864, 75, doi: 10.3847/1538-4357/aad6e0
                                                                   1331 Owen, J. E., & Campos Estrada, B. 2020, Monthly Notices
1281
1282 Koll, D. D. B., & Cronin, T. W. 2019, The Astrophysical
                                                                         of the Royal Astronomical Society, 491, 5287,
                                                                   1332
      Journal, 881, 120, doi: 10.3847/1538-4357/ab30c4
                                                                         doi: 10.1093/mnras/stz3435
1283
                                                                   1333
1284 Kopparapu, R. K., Ramirez, R. M., SchottelKotte, J., et al.
                                                                       Owen, J. E., & Wu, Y. 2013, Astrophysical Journal, 775, 1,
                                                                   1334
      2014, The Astrophysical Journal Letters, 787, L29,
                                                                         doi: 10.1088/0004-637X/775/2/105
1285
                                                                   1335
      doi: 10.1088/2041-8205/787/2/L29
                                                                   1336 Patel, B. H., Percivalle, C., Ritson, D. J., Duffy, C. D., &
1286
1287 Kopparapu, R. K., Ramirez, R., Kasting, J. F., et al. 2013,
                                                                         Sutherland, J. D. 2015, Nature Chemistry, 7, 301,
                                                                   1337
      The Astrophysical Journal, 765, 131,
                                                                         doi: 10.1038/nchem.2202
1288
                                                                   1338
      doi: 10.1088/0004-637X/765/2/131
                                                                       Pierrehumbert, R., & Gaidos, E. 2011, The Astrophysical
1289
                                                                   1339
1290 Lichtenberg, T., & Miguel, Y. 2024, Super-Earths and
                                                                         Journal Letters, 734, L13,
                                                                   1340
      Earth-like Exoplanets, arXiv,
                                                                         doi: 10.1088/2041-8205/734/1/L13
1291
                                                                   1341
      doi: 10.48550/arXiv.2405.04057
                                                                       Quanz, S. P., Ottiger, M., Fontanet, E., et al. 2022,
1292
    Lineweaver, C. H., & Davis, T. M. 2002, Astrobiology, 2,
                                                                         Astronomy & Astrophysics, 664, A21,
                                                                   1343
1293
      293, doi: 10.1089/153110702762027871
                                                                         doi: 10.1051/0004-6361/202140366
   Lingam, M., Nichols, R., & Balbi, A. 2024, Astrobiology,
                                                                       Ramirez, R. M. 2018, Geosciences, 8, 280,
1295
      24, 813, doi: 10.1089/ast.2024.0037
                                                                         doi: 10.3390/geosciences8080280
1297 Luger, R., & Barnes, R. 2015, Astrobiology, 15, 119,
                                                                         -. 2020, Monthly Notices of the Royal Astronomical
                                                                         Society, 494, 259, doi: 10.1093/mnras/staa603
      doi: 10.1089/ast.2014.1231
                                                                   1348
1299 Luque, R., Coy, B. P., Xue, Q., et al. 2024, A Dark, Bare
                                                                   1349 Ramirez, R. M., & Kaltenegger, L. 2014, The Astrophysical
      Rock for TOI-1685 b from a JWST NIRSpec G395H
                                                                         Journal Letters, 797, L25,
                                                                   1350
1300
      Phase Curve, arXiv, doi: 10.48550/arXiv.2412.03411
                                                                         doi: 10.1088/2041-8205/797/2/L25
                                                                   1351
1301
1302 Malaterre, C., Jeancolas, C., & Nghe, P. 2022,
                                                                         -. 2017, The Astrophysical Journal Letters, 837, L4,
                                                                   1352
      Astrobiology, 22, 851, doi: 10.1089/ast.2021.0162
                                                                         doi: 10.3847/2041-8213/aa60c8
                                                                   1353
1303
1304 Mamajek, E., & Stapelfeldt, K. 2023, 55, 116.07
                                                                         -. 2018, The Astrophysical Journal, 858, 72,
                                                                   1354
   Mann, H. B., & Whitney, D. R. 1947, The Annals of
                                                                         doi: 10.3847/1538-4357/aab8fa
                                                                   1355
1305
      Mathematical Statistics, 18, 50
                                                                   1356 Ranjan, S., Nayak, P. K., Pineda, J. S., & Narang, M. 2023,
1306
1307 Melosh, H. J. 2011, Planetary Surface Processes,
                                                                         The Astronomical Journal, 166, 70,
                                                                   1357
      Cambridge Planetary Science (Cambridge: Cambridge
                                                                         doi: 10.3847/1538-3881/ace32d
                                                                   1358
1308
      University Press), doi: 10.1017/CBO9780511977848
                                                                       Ranjan, S., & Sasselov, D. D. 2016, Astrobiology, 16, 68,
                                                                   1359
1309
                                                                         doi: 10.1089/ast.2015.1359
1310 Mojzsis, S. J., Harrison, T. M., & Pidgeon, R. T. 2001,
                                                                   1360
      Nature, 409, 178, doi: 10.1038/35051557
                                                                       —. 2017, Astrobiology, 17, 169, doi: 10.1089/ast.2016.1519
1311
1312 Mol Lous, M., Helled, R., & Mordasini, C. 2022, Nature
                                                                   1362 Ranjan, S., Wordsworth, R., & Sasselov, D. D. 2017a,
      Astronomy, 6, 819, doi: 10.1038/s41550-022-01699-8
                                                                         Astrobiology, 17, 687, doi: 10.1089/ast.2016.1596
1314 Mordasini, C. 2020, Astronomy and Astrophysics, 638, 1,
                                                                        —. 2017b, The Astrophysical Journal, 843, 110,
      doi: 10.1051/0004-6361/201935541
                                                                         doi: 10.3847/1538-4357/aa773e
```

```
1366 Rapf, R. J., & Vaida, V. 2016, Physical Chemistry
```

- 1367 Chemical Physics, 18, 20067, doi: 10.1039/C6CP00980H
- 1368 Revol, A., Bolmont, É., Sastre, M., et al. 2024, Astronomy
- 1369 & Astrophysics, 691, L3,
- doi: 10.1051/0004-6361/202451532
- 1371 Richey-Yowell, T., Shkolnik, E. L., Schneider, A. C., et al.
- 2023, The Astrophysical Journal, 951, 44,
- doi: 10.3847/1538-4357/acd2dc
- 1374 Rimmer, P. B. 2023, in Conflicting Models for the Origin of
- 1375 Life (John Wiley & Sons, Ltd), 407–424,
- doi: 10.1002/9781119555568.ch16
- 1377 Rimmer, P. B., Ranjan, S., & Rugheimer, S. 2021a,
- Elements, 17, 265, doi: 10.2138/gselements.17.4.265
- 1379 Rimmer, P. B., Thompson, S. J., Xu, J., et al. 2021b,
- 1380 Astrobiology, 21, 1099, doi: 10.1089/ast.2020.2335
- 1381 Rimmer, P. B., Xu, J., Thompson, S. J., et al. 2018, Science
- Advances, 4, eaar3302, doi: 10.1126/sciadv.aar3302
- 1383 Rogers, J. G., Gupta, A., Owen, J. E., & Schlichting, H. E.
- 1384 2021, 17, 1
- 1385 Rogers, J. G., & Owen, J. E. 2020, 18, 1
- 1386 Rugheimer, S., Segura, A., Kaltenegger, L., & Sasselov, D.
- 2015, The Astrophysical Journal, 806, 137,
- doi: 10.1088/0004-637X/806/1/137
- Sasselov, D. D., Grotzinger, J. P., & Sutherland, J. D. 2020,
- Science Advances, 6, eaax3419,
- doi: 10.1126/sciadv.aax3419
- 1392 Schlecker, M., Apai, D., Lichtenberg, T., et al. 2024, The
- Planetary Science Journal, 5, 3, doi: 10.3847/PSJ/acf57f
- 1394 Schlecker, M., Pham, D., Burn, R., et al. 2021, Astronomy
- 1395 & Astrophysics, 656, A73,
- doi: 10.1051/0004-6361/202140551
- 1397 Schlichting, H., Sari, R., & Yalinewich, A. 2014, eprint
- arXiv:1406.6435, 1. https://arxiv.org/abs/1406.6435
- 1399 Seager, S., Bains, W., & Hu, R. 2013, The Astrophysical
- Journal, 777, 95, doi: 10.1088/0004-637X/777/2/95
- 1401 Segura, A., Kasting, J. F., Meadows, V., et al. 2005,
- 1402 Astrobiology, 5, 706, doi: 10.1089/ast.2005.5.706
- 1403 Smart, R. L., Sarro, L. M., Rybizki, J., et al. 2021,
- Astronomy & Astrophysics, 649, A6,
- и doi: 10.1051/0004-6361/202039498

- 1406 Spiegel, D. S., & Turner, E. L. 2012, Proceedings of the
- National Academy of Sciences, 109, 395,
- doi: 10.1073/pnas.1111694108
- 1409 Spinelli, R., Borsa, F., Ghirlanda, G., Ghisellini, G., &
- Haardt, F. 2023, The Ultraviolet Habitable Zone of
- 1411 Exoplanets, doi: 10.48550/arXiv.2303.16229
- 1412 Tuchow, N., Stark, C., & Mamajek, E. 2024, HPIC: The
- 1413 Habitable Worlds Observatory Preliminary Input
- 1414 Catalog, arXiv, doi: 10.48550/arXiv.2402.08038
- 1415 Tuchow, N. W., & Wright, J. T. 2023, The Astrophysical
- Journal, 944, 71, doi: 10.3847/1538-4357/acb054
- 1417 Turbet, M., Fauchez, T. J., Leconte, J., et al. 2023, Water
- 1418 Condensation Zones around Main Sequence Stars,
- 1419 https://arxiv.org/abs/2308.15110v1
- 1420 Underwood, D. R., Jones, B. W., & Sleep, P. N. 2003,
- 1421 International Journal of Astrobiology, 2, 289,
- doi: 10.1017/S1473550404001715
- 1423 Vaughan, S. R., Birkby, J. L., Thatte, N., et al. 2024,
- MNRAS, 528, 3509, doi: 10.1093/mnras/stae242
- 1425 Virtanen, P., Gommers, R., Oliphant, T. E., et al. 2020,
- Nature Methods, 17, 261, doi: 10.1038/s41592-019-0686-2
- 1427 Walker, S. I., Bains, W., Cronin, L., et al. 2018,
- 1428 Astrobiology, 18, 779, doi: 10.1089/ast.2017.1738
- 1429 Wang, J., Mawet, D., Ruane, G., Hu, R., & Benneke, B.
- 2017, The Astronomical Journal, 153, 183,
- doi: 10.3847/1538-3881/aa6474
- 1432 West, A. A., Hawley, S. L., Bochanski, J. J., et al. 2008.
- The Astronomical Journal, 135, 785.
- doi: 10.1088/0004-6256/135/3/785
- 1435 Westall, F., Hickman-Lewis, K., Hinman, N., et al. 2018,
- 1436 Astrobiology, 18, 259, doi: 10.1089/ast.2017.1680
- 1437 Zahnle, K. J., & Catling, D. C. 2017, The Astrophysical
- Journal, 843, 122, doi: 10.3847/1538-4357/aa7846
- 1439 Zhang, H., Wang, J., & Plummer, M. K. 2024, AJ, 167, 37,
- doi: 10.3847/1538-3881/ad109e

1459

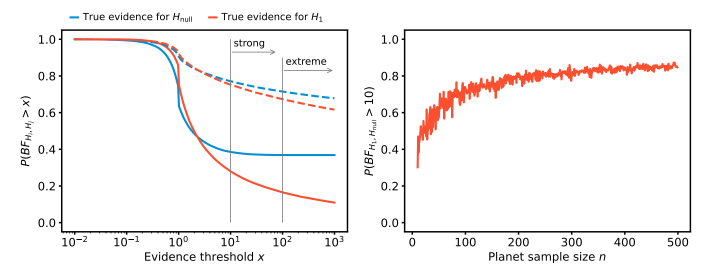


Figure 10. As Figure 3, but using a uniform prior for the abiogenesis rate $f_{\rm life}$. The probabilities of obtaining strong evidence for or against the UV Threshold Hypothesis are overall higher than for the log-uniform prior.

B. ALTERNATIVE PRIORS FOR THE ABIOGENESIS RATE

In the main text, we used a log-uniform prior for the probability of life emerging and persisting f_{life} 1442 in order to reflect our ignorance about this parameter and its order of magnitude. We consider this reasonable choice, as it is agnostic about the scale of f_{life} and assigns equal prior probability to all orders of magnitude. Nevertheless, we repeated our semi-analytical analysis for the more optimistic choice of a prior that is uniform in f_{life} . Figure 10 shows the resulting probabilities of obtaining strong 1447 evidence for or against the UV Threshold Hypothesis under this prior. While the resulting trends are 1448 qualitatively similar, the uniform prior yields overall higher probabilities of such a conclusive test for 1449 all sample sizes.

C. IMPACT OF A LONGER ORIGINS TIMESCALE

While evidence from Earth's history suggests that life may have emerged early (Mojzsis et al. 2001; 1452 Dodd et al. 2017), there is no strong empirical basis to assume that this timescale is representative of 1453 habitable planets in the Universe. The emergence of life could be a slow and rare process, possibly requiring much longer periods (Lineweaver & Davis 2002; Spiegel & Turner 2012).

To assess the implications of a more conservative abiogenesis timescale, we consider a scenario in 1455 which a planet's minimum time required to be both in the HZ and having above-threshold NUV fluxes before life can emerge is increased to $\Delta T_{\min} = 100$ Myr, rather than 1 Myr as assumed above. Figures 11, 12, and 13 show the results of our Bioverse simulations under this assumption.

The overall impact differs significantly between M dwarf and FGK dwarf planets. While M dwarf 1460 planets remain largely unaffected, the fraction of inhabited FGK dwarf planets decreases sharply due to the limited overlap between the time of high UV flux and habitable conditions (compare Figure 2). Less the optimistic assumption that the probability of abiogenesis is unity ($f_{\text{life}} = 1$), the fraction of inhabited planets under the UV Threshold Hypothesis declines rapidly with increasing UV flux thresholds (Figure 11).

The impact of a longer origins timescale on testing the UV Threshold Hypothesis becomes evident when we repeat our example survey ($f_{\text{life}} = 0.8$, $F_{\text{NUV,min}} = 300 \, \text{erg s}^{-1} \, \text{cm}^{-2}$) under this assumption (Figure 12): While the survey of M dwarf planets remains largely unaffected, the fraction of inhabited FGK planets is reduced to zero, making a conclusive test of the UV Threshold Hypothesis impossible. Figure 13 demonstrates that this result is independent of the threshold NUV flux and the intrinsic 1473 abiogenesis rate. Given the lack of constraints on the timescale of abiogenesis, these findings highlight the advantages of focusing on M dwarf planets when searching for biosignatures. If life emerges slowly, 1475 FGK planets may rarely, if ever, reach the inhabited stage, whereas M dwarf planets remain viable

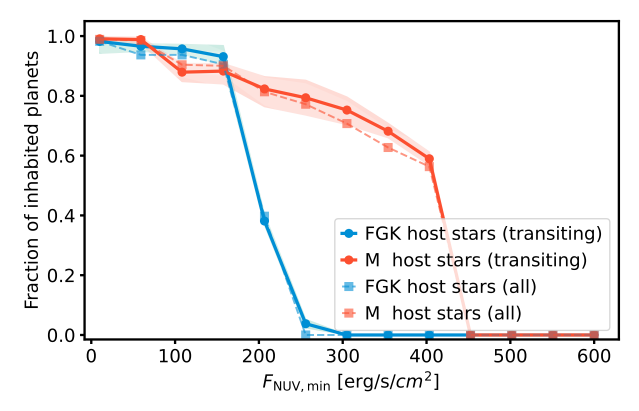


Figure 11. Fraction of inhabited planets as a function of NUV threshold flux for $\Delta T_{\rm min}=100$ Myr. The decrease in inhabited planets is more pronounced for FGK stars, narrowing the viable range of UV Thresholds compared to shorter origins timescales.

targets for future surveys. If biosignatures were nevertheless detected on FGK planets despite the the constraints imposed by a longer abiogenesis timescale, it would suggest that either the UV Thresh-1478 old Hypothesis, as formulated here, is incorrect or that the timescale of abiogenesis is shorter than $\mathcal{O}(100\,\mathrm{Myr})$.

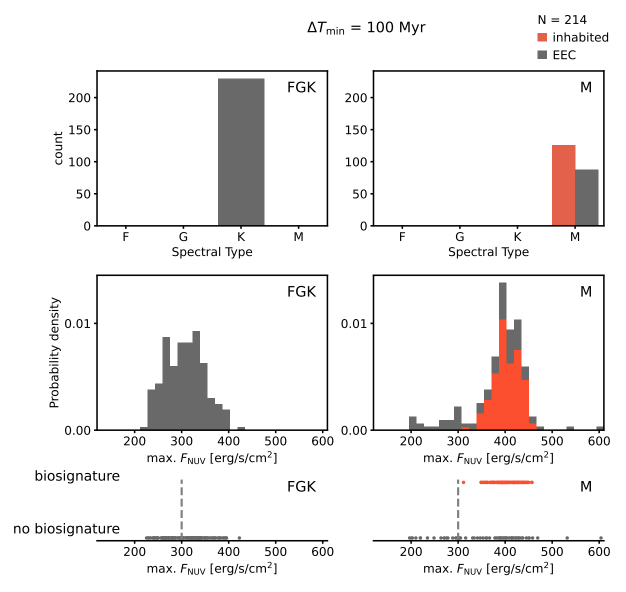


Figure 12. Simulated survey results under the assumption of $\Delta T_{\rm min} = 100$ Myr. The fraction of inhabited FGK planets is reduced to zero, while the fraction of inhabited M dwarf planets remains largely unaffected.

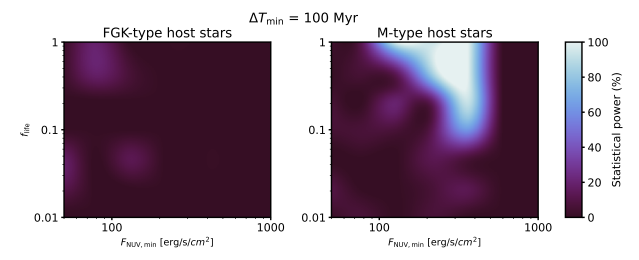


Figure 13. Statistical power for different abiogenesis rates and NUV threshold fluxes under the assumption of $\Delta T_{\rm min}=100$ Myr. The longer origins timescale makes a test of the UV Threshold Hypothesis impossible for FGK planets, while M dwarf planets remain viable targets.