Manuscript version: Author’s Accepted Manuscript
The version presented in WRAP is the author’s accepted manuscript and may differ from the published version or Version of Record.

Persistent WRAP URL:
http://wrap.warwick.ac.uk/98794

How to cite:
Please refer to published version for the most recent bibliographic citation information. If a published version is known of, the repository item page linked to above, will contain details on accessing it.

Copyright and reuse:
The Warwick Research Archive Portal (WRAP) makes this work by researchers of the University of Warwick available open access under the following conditions.

Copyright © and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable the material made available in WRAP has been checked for eligibility before being made available.

Copies of full items can be used for personal research or study, educational, or not-for-profit purposes without prior permission or charge. Provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.

Publisher’s statement:
Please refer to the repository item page, publisher’s statement section, for further information.

For more information, please contact the WRAP Team at: wrap@warwick.ac.uk.
DYNAMICAL AND BIOLOGICAL PANSPERMIA CONSTRAINTS WITHIN MULTI-PLANET EXOSYSTEMS

Dimitri Veras†
Centre for Exoplanets and Habitability, University of Warwick, Coventry, CV4 7AL, UK and
Department of Physics, University of Warwick, Coventry, CV4 7AL, UK

David J. Armstrong
Centre for Exoplanets and Habitability, University of Warwick, Coventry, CV4 7AL, UK and
Department of Physics, University of Warwick, Coventry, CV4 7AL, UK

James A. Blake
Centre for Exoplanets and Habitability, University of Warwick, Coventry, CV4 7AL, UK and
Department of Physics, University of Warwick, Coventry, CV4 7AL, UK

Jose F. Gutiérrez-Marcos
Centre for Exoplanets and Habitability, University of Warwick, Coventry, CV4 7AL, UK and
School of Life Sciences, University of Warwick, Coventry, CV4 7AL, UK

Alan P. Jackson
Centre for Planetary Sciences, University of Toronto at Scarborough, Toronto, ON, Canada and
School of Earth and Space Exploration, Arizona State University, AZ, USA

Hendrik Schäfer
Centre for Exoplanets and Habitability, University of Warwick, Coventry, CV4 7AL, UK and
School of Life Sciences, University of Warwick, Coventry, CV4 7AL, UK

ABSTRACT

As discoveries of multiple planets in the habitable zone of their parent star mount, developing analytical techniques to quantify extrasolar intra-system panspermia will become increasingly important. Here, we provide user-friendly prescriptions that describe the asteroid impact characteristics which would be necessary to transport life both inwards and outwards within these systems within a single framework. Our focus is on projectile generation and delivery and our expressions are algebraic, eliminating the need for the solution of differential equations. We derive a probability distribution function for life-bearing debris to reach a planetary orbit, and describe the survival of micro-organisms during planetary ejection, their journey through interplanetary space, and atmospheric entry.

Subject headings: panspermia; extrasolar terrestrial planets; habitable zone; impact processes; planetary habitability and biosignatures

1. INTRODUCTION

Although studies of the transport of life-bearing rocks between planets have a long history (e.g. Melosh [1988], claims of the discovery of traces of ancient life in the meteorite ALH84001 in the mid-1990s [McKay et al. 1996] accelerated investigations into panspermia within the Solar system. The last two decades have since featured detailed work (e.g. Mileikowsky et al. 2000; [Mileikowsky et al. 2000]) outlining delivery dynamics [Melosh & Tonks 1993; Gladman et al. 1996; Gladman 1997; Alvarellos et al. 2002; Reves-Ruiz et al. 2012; Worth et al. 2013], impact physics and chemistry [Meyer et al. 2011; Rubin 2015; Barney et al. 2016], and biological survival requirements [Horneck et al. 1994; Mastrapa et al. 2001; Nicholson 2009; Moeller et al. 2010] with respect to Earth, Mars and other solar system bodies. Consequently, a detailed foundation for panspermia-related processes has been established.

Despite these advances, the applicability of these processes to extrasolar planetary systems is still in question, partly because in those systems we lack the detailed knowledge of our own planetary system. Nevertheless, efforts to characterize panspermia between different extrasolar systems, or between the solar system and extrasolar systems, have contributed to our understanding (Adams & Spergel 2005; Valtonen et al. 2009; Wesson 2010; Belbruno et al. 2012; Lin & Loeb 2013; Gale & Wandel 2013). However, panspermia amongst extrasolar planets within the same system has received a relatively little but increasing amount of attention (Heller & Armstrong 2014; Steffen & L 2016; Krijt et al. 2017; Lingam & Loeb 2017a). A potential reason for this relative dearth of studies is the lack of observational evidence of multiple planets in the habitable zone of the same star.
This situation has now changed with the ground-breaking discovery of multiple potentially habitable planets in the TRAPPIST-1 system (Gillon et al. 2017). TRAPPIST-1 is an M dwarf with a mass of 0.08 M☉ that harbours seven observed transiting planets all with masses similar to the Earth and three (planets e, f and g) which are securely in the star’s habitable zone (although all seven may be, with effective temperatures ranging from about 150 K to 400 K). Because all seven planets are seen transiting from the Earth, their orbits are nearly coplanar. The system is compact (all planets could fit well within Mercury’s orbit), and are likely to be resonantly interacting in a long chain (Lager et al. 2017). We do not yet know if an equivalent of the Late Heavy Bombardment event (believed to have occurred in the Solar system, even if less intense than originally thought; Bottke & Norman 2017) has or will occur in that system, nor what types of potential impactors lurk beyond the most distant planet (h) and outside of our field of view. Despite the uncertainties, investigation of panspermia from within this system has already been undertaken (Krijt et al. 2013; Lingam & Loeb 2017a) and might be prompted further by additional observations, which are currently underway.

Here, we study and derive several aspects of lithopanspermia in more general closely-packed multi-planet systems, with a focus on analytics and dynamical delivery, but also addressing micro-organism survival at each stage. Numerous Solar system studies have taught us that N-body simulations are both computationally expensive and dependent on a large number of parameters (Dones et al. 1999) which are unknown in exoplanetary systems. Computational times for most known extrasolar planetary systems would be worse because of their compact nature. Therefore, we adopt a purely analytical approach, one which could be applied to extrasolar systems with multiple habitable planets. The characterisation of such systems is expected to increase steadily over the next decade, culminating with the PLATO mission (Rauer et al. 2014), which will measure habitable planets out to about 1 au.

Throughout the paper, our subscript convention for physical quantities will be: $i$ for the impactor, no subscript for the source planet, a single prime for the fragmented debris ejected from the source planet, and a double prime for the target planet. In Section 2, we establish our setup and describe how a life-bearing rock could be transferred between one planet and the orbit of another planet: Appendix A provides most of the intermediate equations required for this section, and Appendix B contains an extension with a fictitious template compact system which can be used for quick estimates. Section 3 then details the likelihood of that rock actually impacting a target planet. Section 4 constrains the characteristics of the ejecta that would both satisfy the dynamics and have the capability to harbour life. In Section 5, we consider the biological prospects of life surviving all aspects of lithopanspermia. We conclude in Section 6.

2. ORBIT TRANSFERS

2.1. Setup

Consider a pair of planets such that one, the “source”, contains a life-bearing organism, whereas the other, the “target”, initially does not. An impactor crashes into the source, producing a spray of life-bearing ejecta. By assuming that the ejecta is “kicked” impulsively, we estimate its direction and speed such that it would reach the orbit of the target. By impulsively, we refer to the timescale of the kick being much smaller than the orbital period of the source. In multi-planet systems within the detectability threshold of transit photometry surveys, planet orbital periods are on the order of days, whereas impact kick timescales would typically be on the order of minutes.

The underlying formalism we use was established in Jackson & Wyatt (2012) and expanded upon in Jackson et al. (2014), and similar to that in Appendix A of Le Feuvre & Wieczorek (2008). We briefly repeat here the geometrical setup in Jackson et al. (2014): Denote the kick speed as $\Delta v$, and the circular speed of the source as $v_k \equiv \sqrt{G (M_{\text{star}} + M) / a}$, where $a$ and $M$ are the source’s semimajor axis and mass. The launch speed $v_{\text{lau}}$ and escape speed $v_{\text{esc}}$ are related to $\Delta v$ through $v_{\text{lau}}^2 = (\Delta v)^2 + v_{\text{esc}}^2$, where $v_{\text{esc}}^2 = 2GM/R$.

For perspective, the circular speed of the TRAPPIST-1 planets are, from planet b to h moving outward, $\{79.9, 68.2, 57.5, 50.2, 43.7, 39.6, 33.9\}$ km/s. The ratio $\Delta v / v_k$ features frequently in the equations, and is bound from above in our study by the value of $\sqrt{2} + 1$, which is the maximum possible value for which the debris can remain in the planetary system. Further, the minimum value of $\Delta v / v_k$ for which the ejecta can escape the system is $\sqrt{2} - 1$.

At impact, the source is assumed to lie at the pericentre of its orbit such that its argument of pericentre, longitude of ascending node, and inclination $I$ are all zero. The source is assumed to move counterclockwise, and the kick direction is defined by two variables: $\theta$ and $\phi$. The angle between the source’s angular momentum vector and the kick direction is $\theta$ and the angle between the star-source pericentre line and the projection of the kick direction onto the source’s orbital plane is $\phi$.

2.2. Ejecta orbit characteristics

The orbit of the ejecta, whose elements are denoted by primes, are related to the unprimed quantities (which refer to the source planet), through equations (A1)-(A8) (Jackson et al. 2014). Because known compact multi-planet systems are dynamically “cold” – exhibiting low eccentricities – henceforth we make the approximation that all planets are on circular orbits. Doing so greatly simplifies the analysis. For example, the upper limits for the planetary eccentricities in the TRAPPIST-1 system are all under 0.085. Further, because for any compact multi-planet system with habitable plants, we cannot yet know if panspermia has occurred, or will occur, at a particular time, we assume, without loss of generality, that the true anomaly $f = 0$. We hence constrain only when ejecta intersects the orbit of the target, and use equations (A7) and (A8) for this purpose.

2.3. Coplanarity restrictions

$^1$ The restrictiveness of the assumption of impact at orbital pericentre will be removed later when we assume circular orbits.
First however, consider that in order for ejecta to hit the target, the ejecta must coincide with the target in all three spatial dimensions. This intersection is most easily achieved if the orbital planes of the source, ejecta and target lie close to one another as we can then be guaranteed of coincidence in one of the three dimensions. Hence, in this section, we quantify how coplanar the orbits of the ejecta, source and target must be to achieve spatial coincidence in the vertical direction (direction of the angular momentum vector of the source) throughout the debris orbit.

This condition is mathematically equivalent to \( q \sin I' \lesssim R \) or \( Q' \sin I' \lesssim R \), when assuming that \( R \) denotes planet radius, \( q' \) denotes orbital pericentre, \( Q' \) denotes orbital apocentre, and the reference plane is the one which connects a coplanar source and target. Note that unless the longitudes of ascending node can be measured, the mutual inclinations of all source-target pairs in a particular system will remain unknown. The approximations in these relations result from effects not considered here such as gravitational focusing and atmospheric drag.

If the kick direction is perpendicular to the source’s angular momentum direction (\( \theta = \pi/2 \)), then the ejecta orbit will be coplanar with the reference orbit (\( I' = 0 \)). If, however, the kick direction deviates from these values, then the result is less obvious and is dependent on \( \phi \) (and \( e \) when not assumed to be zero).

Figure 1 illustrates the resulting dependence of \( I' \) on \( \phi \) for two different values of \( \theta \), one where \( \theta \) deviates from coplanarity by \( 10^{-3} \) degrees (left panel) and the other where the deviation is \( 0.5 \) degrees (right panel).

In order to provide some context, we superimpose some results from the TRAPPIST-1 system on this figure. The gray region corresponds to where in the TRAPPIST-1 system \( I' \leq \arcsin (\min (a/R)) = 0.029^\circ \) and the red region to where \( I' \geq \arcsin (\max (a/R)) = 0.24^\circ \). For this red region, the ejecta and target would be vertically coincident only near the nodes of their orbits, whereas for the gray region, they would be vertically coincident throughout their orbits (increasing the chances of impact). The purple curve, corresponding to \( \Delta v/v_k = \sqrt{2} + 1 \), is a limiting case for which the ejecta may remain bound. As \( \Delta v/v_k \) is lowered and approaches zero, the resulting curves would be lower than the green curve. For a kick deviation offset from coplanarity exceeding about \( 0.86^\circ \), we find that all three curves lie entirely within the red region.

2 In the TRAPPIST-1 system, the longitudes of ascending node are so far unknown, and the maximum difference in measured inclinations is \( 0.21^\circ \), when neglecting errors.

### 2.4. Other spatial restrictions

Having linked the angles of impact with the inclinations, we now turn to the other two spatial dimensions. The subsequent analysis greatly benefits from three simplifications, which are sufficient for this study. The first two are our continued assumption of circular and coplanar orbits. The third is that we consider the source and target planets in pairs, ignoring the influence of any other planets including those whose orbits lie in-between the pairs. This last assumption degrades in accuracy as the distance between the planets increases, because the debris will take longer to traverse this distance, and hence be diverted to a greater extent by extra bodies. However, these effects are sensitively dependent on the number, masses and locations of other planets in the system.

#### 2.4.1. Intersecting orbits

In order for a collision to occur, the debris must be ejected onto an orbit which intersects the orbit of another planet. We can place bounds on this geometry by considering the pericentre and apocentre. Hence we begin by expressing these quantities as a function of the impacted planet’s semimajor axis, the velocity ratio and \( \phi \) (equations A9-A11). We wish to find the range of velocity kicks for which these pericentres or apocentres are achieved. Hence, inverting the equations yields, for in-
ward motion towards the orbital pericentre,
\[
\left( \frac{\Delta v}{v_k} \right)_{\text{in}} = \begin{cases} 
\frac{(a-q')(a+q')}{q'^2 - a^2 \sin^2 \phi} \sin \phi + \sqrt{\zeta}, & \sin^2 \phi < \left( \frac{q'}{a} \right)^2 \\
\frac{(a-q')(a+q')}{q'^2 - a^2 \sin^2 \phi} \sin \phi - \sqrt{\zeta}, & \sin^2 \phi > \left( \frac{q'}{a} \right)^2
\end{cases}
\]  
(1)
where
\[
\zeta = q' \left[ q' + (2a + q') \sin^2 \phi \right].
\]  
(2)
Recall that \( a \) and \( q' \) refer respectively to the source planet’s semimajor axis and the ejecta’s orbital pericentre, and \( \phi \) is the angle between the projection of the kick direction onto the source’s orbital plane and the star-source pericentre line.

For panspermia outward from the star, no piecewise function is necessary, as in the following equation the denominator is always positive and the term in square brackets is always negative.
\[
\left( \frac{\Delta v}{v_k} \right)_{\text{out}} = \frac{(a - Q') \left[ (a + Q') \sin \phi - \sqrt{\zeta} \right]}{Q'^2 - a^2 \sin^2 \phi}
\]  
(3)
where
\[
\zeta = Q' \left[ Q' + (2a + Q') \sin^2 \phi \right].
\]  
(4)
Equations (1)-(4) provide the necessary and sufficient conditions for kick speeds to propel ejecta into the orbit of another planet.

2.4.2. Application to TRAPPIST-1

In order to demonstrate how these equations can be applied to a real system, Fig. 2 illustrates the application of equation (1) to the TRAPPIST-1 system. The figure displays inward panspermia from planet h, as well as what minimum kick speeds are necessary to thrust the ejecta just into the orbits of planets g, f, e, d, c and b. Note that this value is highly dependent on \( \phi \), and exceeds the system escape speed for many values of \( \phi \). Further, the dependence on \( \phi \) is non-monotonic.

When we instead consider outward panspermia to planet h, the functional form changes. Application of equation (3) yields Fig. 3. Here, for all values of \( \phi \), the minimum kick speed required to propel the ejecta to another planet’s orbit is smaller than the system escape speed. The smallest kick would be from Planet h’s nearest neighbour (planet g), whereas the greatest kick is required from the planet furthest away (planet b).

Both Figs. 2 and 3 do not take into account the escape speed from the planets themselves. The escape speed of the TRAPPIST-1 planets are, from planet b to h moving outward, \( \{9.90, 12.79, 8.15, 9.19, 9.02, 12.2, 9.35\} \) km/s. The relative velocity of the ejecta after escape from the planet is likely to be comparable to the escape velocity of the planet.

2.4.3. When keeping \( \phi \) fixed

If instead one has reason to assume a particular fixed value of \( \phi \), then equations (1)-(4) may be considered as functions of \( q'/a \) or \( a/Q' \). As an example, the result for outward panspermia is shown in Fig. 4 for six values of \( \phi \). This plot may be applied to any planetary system, including the solar system. For example, outward panspermia from Earth to Mars (Worth et al. 2013) corresponds roughly to the right axis of the plot. The escape speeds of those planets, however, differ by over an order of magnitude from those of the TRAPPIST-1 planets.

2.4.4. Curve extrema

Returning to Figs. 2 and 3, in order to find the minimum value of these curves as a function of \( \phi \), we consider all possible extrema of equation (1), which are found in equation (A12). For outward panspermia,
\[
\max \left( \frac{\Delta v}{v_k} \right)_{\text{out}} = \sqrt{\frac{2Q'}{a + Q'} + 1}
\]  
(5)
Outward panspermia as a function of \((a/Q')\) when \(\phi\) is kept fixed. The x-axis includes the entire relevant range in the TRAPPIST-1 system; the upper end also roughly corresponds to outward panspermia from Earth’s orbit to Mars’ (assuming that they are on circular, coplanar orbits).

and \((\phi)_{\text{ex}} = \pi/2\), which gives the absolute minimum

\[
\min \left( \frac{\Delta v}{v_k} \right)_{\text{out}}^{\text{abs}} = \sqrt{2Q'} - 1. \tag{6}
\]

These equations explain why the system escape speed corresponding to \(\Delta v/v_k (= \sqrt{2} + 1)\) is never reached. Because the edge of the system is beyond planet h, the ejecta will reach planet h before the edge of the system.

Alternatively, for inward panspermia, there are multiple real solutions: \((\phi)_{\text{ex}} = \{ -\pi/2, -\arccos \left( \pm \sqrt{a^2 - q'^2}/a \right) \}\). Combined with the escape boundary, we have

\[
\max \left( \frac{\Delta v}{v_k} \right)_{\text{in}}^{\text{abs}} = \sqrt{2} + 1, \tag{7}
\]

\[
\min \left( \frac{\Delta v}{v_k} \right)_{\text{in},1} = \sqrt{\frac{2q'}{a + q'}} + 1, \tag{8}
\]

\[
\min \left( \frac{\Delta v}{v_k} \right)_{\text{in},2} = \frac{a(a-q')}{2q'(a+q')}. \tag{9}
\]

The absolute minimum could arise from either equations (6) or (7). Equation (6) is the absolute minimum when \(q'/a < (1/2)(\sqrt{2} - 1) \approx 0.21\); otherwise equation (7) gives the absolute minimum. Equations (6)-(9) may be useful constraints for any compact multi-planet system.

2.4.5. Probability distributions

Given these constraints, we can now construct probability distribution functions for debris reaching the target planet’s orbit. We obtain a probability \(P\) of a given ratio \((\Delta v/v_k)\) for either \(\phi\) either 

\[
\min \left( \frac{\Delta v}{v_k} \right)_{\text{out}}^{\text{abs}} = \sqrt{2Q'} - 1. \tag{6}
\]

\[
\max \left( \frac{\Delta v}{v_k} \right)_{\text{in}}^{\text{abs}} = \sqrt{2} + 1, \tag{7}
\]

\[
\min \left( \frac{\Delta v}{v_k} \right)_{\text{in},1} = \sqrt{\frac{2q'}{a + q'}} + 1, \tag{8}
\]

\[
\min \left( \frac{\Delta v}{v_k} \right)_{\text{in},2} = \frac{a(a-q')}{2q'(a+q')}. \tag{9}
\]

The absolute minimum could arise from either equations (6) or (7). Equation (6) is the absolute minimum when \(q'/a < (1/2)(\sqrt{2} - 1) \approx 0.21\); otherwise equation (7) gives the absolute minimum. Equations (6)-(9) may be useful constraints for any compact multi-planet system.

2.4.5. Probability distributions

Given these constraints, we can now construct probability distribution functions for debris reaching the target planet’s orbit. We obtain a probability \(P\) of a given ratio \((\Delta v/v_k)\) being sufficiently high for the ejecta to reach the orbit of another planet as an explicit function of \((\Delta v/v_k), a\) and either \(q'\) or \(Q'\), by assuming some distribution for \(\phi\). Because the cratering record on Solar system bodies indicates that ejecta are effectively isotropically distributed, we assume that a uniform distribution of \(\phi\) holds generally for other planetary systems. Consequently, the formulae we obtain may be applied widely if extrasolar systems experience similar cratering processes. Our final results are presented in equations (A18)-(A24).

Now we provide an example using these probability functions. We utilize the TRAPPIST-1 system both in order to be consistent with the previous applications and because, fortuitously, planets b and h are sufficiently well-separated to allow us to sample a special case where \(q'/a < (1/2)(\sqrt{2} - 1)\). Recall that the only variables required to construct these functions are the semimajor axes of the planets and the velocity ratio.

Figures 5 and 6 are the result. Note first that for inward panspermia, \(P\) never reaches unity, unlike for outward panspermia: the geometry responsible for these relations are folded into the curves. Second, for both directions, transferring to neighbouring planets is easier than for those further away. Third, the kinks in Fig. 5ulti-
3. IMPACTING THE TARGET IN ONE PASS

In the last section, we placed bounds on orbital properties of ejecta that could hit the target planet. In order to better quantify the probability of actually impacting the target, we now consider the location of the debris in space, rather than just its orbit.

Assuming that the ejecta orbit intersects or almost intersects with the orbit of the target, and that these orbits remain fixed, then expressions exist for the probability of collision in one pass. These expressions, pioneered by Opik (1951) and Wetherill (1967), have led to substantial and wide-ranging applications. A recent updated and simplified series of derivations was provided by JeongAhn & Malhotra (2017). They found in their equations 29 and 37 the probability of collision per revolution of the ejecta, \( P \), as a function of (i) the orbital periods of the target and ejecta, (ii) the gravitational acceleration due to the parent star (assumed to be constant in the vicinity of the collision), (iii) the collision radius, (iv) the velocities of both objects at collision, and (v) the angle, \( \lambda \), between the common direction of the velocity vectors and the vector from the star to the collision point.

We now consider these dependencies in more detail. Because we assume that the target is on a circular orbit, the gravitational acceleration at the collision point is \( GM_{\text{star}}/v^{\prime \prime 2} \) (recall that the target’s orbital parameters and mass are denoted with double primes). The collision radius is the sum of the radii of the ejecta and target multiplied by the gravitational focusing factor \( (1 + v^{\prime \prime 2}_{\text{esc}}/v^{\prime \prime 2}) \). These velocities can be expressed in orbital elements as \( v^{\prime \prime} = n^{\prime \prime}a^{\prime \prime} \) and \( v^{\prime} = n^{\prime}a^{\prime} \sqrt{(1 + e^{\prime 2} \cos f^{\prime})/(1 - e^{\prime 2})} \) such that \( n^{\prime} \) and \( n^{\prime \prime} \) denote the mean motion of the ejecta and target. In this last expression, we can further relate \( f^{\prime} \) to \( \phi \). For the circular source orbit case, and assuming coplanarity amongst the source, debris and target, equations (A4) and (A5) reduce to

\[
\tan f^{\prime} = \frac{1 + \left( \frac{\Delta v}{v^{\prime \prime}_{\text{esc}}} \right) \sin \phi}{2 + \left( \frac{\Delta v}{v^{\prime \prime}_{\text{esc}}} \right) \sin \phi} \quad (10)
\]

which leads to Fig. 7.

Adding these various components above together leads to the following expression for the collision probability per time in our formalism. Assuming that the target and ejecta are much less massive than the parent star allows us to concisely write

\[
P = \frac{c}{4\pi^2} \sqrt{G \left( \frac{M_{\text{star}}^2}{M^{\prime \prime}} \right) \left( \frac{R^{\prime} + R^{\prime \prime}}{a^{\prime}a^{\prime \prime}} \right)} \times \frac{\sqrt{|v^{\prime} - v^{\prime \prime}|}}{v^{\prime} + v^{\prime \prime}} \left( 1 + \frac{v^{\prime \prime 2}_{\text{esc}}}{|v^{\prime} - v^{\prime \prime}|^2} \right) \csc \lambda \quad (11)
\]

where \( c \) is a constant equal to either \( \sqrt{2} \) for exactly intersecting orbits, or 1.7 for non-intersecting orbits. Equation (11) accounts for both inward and outward panspermia because of the first absolute value. As emphasized by JeongAhn & Malhotra (2017), equation (11) should not be used to predict a specific impact event in the past or future, but rather be used in a statistical way with streams of debris.

The time taken for the debris to travel from the source to the target planet – a measure which influences the survival rates of microbes – may be approximated by the inverse of equation (11). This transit time is hence strongly influenced by \( |v^{\prime} - v^{\prime \prime}| \), which is the difference in speed of the ejecta and target planet. When these values are comparable, the inverse of equation (11) becomes singular. Further, the angle \( \lambda \) is crucial, and can reduce the transit time almost arbitrarily. These factors primarily explain the several order-of-magnitude spread in transit times in fig. 3 of Krijt et al. (2017) for the TRAPPIST-1 system. Nevertheless, when comparing transit times in TRAPPIST-1 to other systems, the functional dependencies in equation (11) may be useful. For example, the transit time scales inversely with the mass of the star and as the square root of the distance from the target planet.

4. EJECTA CHARACTERISTICS

Alternatively, if we assume that all of the ejecta represents a single boulder or pebble of mass \( m^{\prime} \), then we can speculate on the characteristics of this piece of ejecta, and whether it could be life-bearing. In this section, we neglect more complex possibilities, such as indirect supplementary ejection from small source planets. Source planets the size of Ceres, for example, have low-enough escape velocities and surface pressures to be susceptible to glaciopanspermia (Houtkooper 2011; Lingam & Loeb 2017). We also neglect the portion of the atmosphere – even for large source planets – that will invariably be ejected along with the debris (Berera 2017). The debris can act as an agent to transfer the source planet’s atmospheric constituents to the target. If the transfer of material is extensive enough, as during perhaps a period of heavy bombardment (de Niem et al. 2012; Bottke & Norman 2017), then the atmosphere of the target planet may become less or more susceptible towards hosting life on its surface.
A piece of ejecta which could escape an atmosphere is planet contains an atmosphere with surface pressure to maintain escape velocity from the source. If the source This radius must be larger than the critical radius needed magnitude. Otherwise, most post-impact fragments with approximately at least the same mass as Big Ben (the bell) or the Hubble Space Telescope can escape an exo-Earth.

4.1. Escaping the atmosphere

First, consider that the radius and density of this piece of ejecta are related to \( m' \) through \( m' = \frac{4\pi}{3}\rho R^3 \). This radius must be larger than the critical radius needed to maintain escape velocity from the source. If the source planet contains an atmosphere with surface pressure \( p \) and gravitational acceleration \( g \), then the radius \( R' \) of a piece of ejecta which could escape an atmosphere is [Artemieva & Ivanov 2004]

\[
R' \geq \frac{3p}{8g\rho} \left[ \frac{\Delta v + v_{\text{esc}}}{\Delta v - v_{\text{esc}}} \right], \quad \Delta v > v_{\text{esc}} \tag{12}
\]

where \( g = \frac{GM}{R^2} \) such that any ejector could escape from an atmosphere-less source as long as \( \Delta v \) is sufficiently high. Consequently, the minimum single-body ejecta mass which eventually hits the target and could initially escape from the source is

\[
m' \geq \frac{9\pi}{128} \left( \frac{\rho \rho'}{\rho^2 g^2} \right) \left[ \frac{\Delta v + v_{\text{esc}}}{\Delta v - v_{\text{esc}}} \right]^3, \quad \Delta v > v_{\text{esc}}. \tag{13}
\]

We plot equation (13) in Fig. 8 assuming that the source planet is an Earth-analog. The plot illustrates that atmospheric drag has the strongest effect when the post-impact speed is close to the escape speed, and flattens out as \( \Delta v \) increases (note that for large \( \Delta v \), equation 13 becomes independent of \( \Delta v \)).

4.2. Largest fragment

Recall that \( m' \) represents a fragment of the impact. Now we consider how to compute the largest possible size of an impact fragment, with radius \( R'_{\text{max}} \). This deduction is based on detailed physics (porosity, spalling, Grady-Kipp processes, simple versus complex crater formation) that are beyond the scope of this study. Further, certain dependencies which might work for icy Solar system satellites [Bierhaus et al. 2012; Singer et al. 2013; Alvarellos et al. 2017] might not apply uniformly to all planets in extrasolar systems.

We just use [Mileikowsky et al. 2000]

\[
\frac{R'_{\text{max}}}{R_i} = \left( \frac{3 + W}{2} \right) \left[ \frac{T}{\rho \left( \Delta v^{2/3} / v_i^{4/3} \right)} \right] \tag{14}
\]

where \( R_i \) and \( v_i \) are the radius and impact speed of the impactor, \( T \) is the tension at fracture, and \( W \) is the Weibulls modulus [Afferrante et al. 2006]. For context, as mentioned by Mileikowsky et al. (2000), \( T = 0.1 \times 10^9 \) Pa and \( W = 9.5 \) for basalt, a type of igneous rock.

We may relate the pre-impact and post-impact speed from equation (14) by assuming that the energy of the impact is deposited at a depth which is comparable to \( R_i \). Then [Melosh 1984; 1988]

\[
\frac{\Delta v}{v_i} \approx \left( \frac{R_i}{d} \right)^{2.87} \tag{15}
\]

where \( d > R_i \) is the distance from the ejection point to the centre of energy deposition[3]. Consequently, the ejecta speed cannot be larger than the impactor speed. Otherwise, the speed ratio is largely determined by the detailed physics of the impact. As we do not pursue such detail here, we suffice to leave \( \Delta v/v_i \) as a free parameter ranging from 0 to 1. Then, we can express equation (14) for basalt (adopting \( \rho = 3 \text{ g/cm}^3 \)) as

\[
\frac{R'_{\text{max}}}{R_i} = \left( \frac{v_i}{0.456 \text{ km/s}} \right)^{-2} \left[ \frac{\Delta v}{v_i} \right]^{-2/3} \tag{16}
\]

We plot equation (16) in Fig. 9. The plot illustrates that, in general, fragment radii are no more than about 5% of the radius of their progenitor.

4.3. Liberating material

\[3 \text{ See Housen et al. (1983) and Dobrovolskis et al. (2010) for alternative formulations.} \]
Fig. 10.— The minimum impactor size that could liberate material from the planet’s gravitational well. The impactor and planet have densities of $3 \text{ g/cm}^3$, which is consistent with basalt. The figure illustrates that panspermia could occur typically only if the impactor size is at least on the order of tens of km in size.

Now we may consider the minimum impactor size that could liberate material. If $R_i = R_i^{max}$ is taken to be the minimum size of ejecta that can escape the atmosphere (through equation (12)), then it follows that the corresponding $R_i = R_i^{min}$ (through equation (16)) must be the minimum size impactor that can liberate material. Combining these equations, and assuming basalt for the impactor and an Earth-sized, Earth-mass planet, yields Fig. 10. The figure demonstrates that generally an impactor must have a radius of at least tens of km in order to be the catalyst for panspermia.

4.4. Destroying the source

Finally, if the impactor is too energetic, then it could sterilize, eviscerate or break apart the source planet. Any of these processes would inhibit prospects for panspermia. Here, we do not analyze the consequences in detail, but merely provide bounds for the impactor in the most extreme case of breakup.

We can place an upper bound on the maximum size of the impactor by considering the maximum specific energy $E_{max}$ imparted to a source for which the source will remain intact. The conditions for catastrophic disruption have an extensive associated literature, and can be characterized through a variety of metrics. For example, see Benz & Asphaug (1999) and the hundreds of more recent papers which cite that one.

We adopt the explicit formalism in Section 5 of Movshovitz et al. (2016). They show that the source planet would break apart if the following condition is met:

$$\left( \frac{v_i}{v_{esc}} \right)^2 > \left( 1.10 \pm 0.58 \right) \times \left[ 8 + 3 \left( \frac{R_i}{R} \right)^{-3} - 5 \left( \frac{R_i}{R} \right) + 8 \left( \frac{R_i}{R} \right)^2 + 3 \left( \frac{R_i}{R} \right)^5 \right].$$

(21)

Figure 11 illustrates the phase space of equation (21). The plot demonstrates that for common definitions of asteroid and planet, an asteroid could never destroy a planet. However, the result of a Mars-sized object colliding with an Earth-sized object is less clear.

4.5. Temperature at impact

We now have some idea of the size of potential ejectors. Returning our attention to panspermia, we must also
consider temperature and how it will affect the ability of life to survive such an impact. Fig. 4 of [Weiss & Head 2016] illustrates averaged ejecta temperatures as a function of both crater diameter and surface heat fluxes on Mars. They find a range of 215 - 600 K for crater diameters between 0 - 150 km in roughly linear relationships for surface heat fluxes of 20 - 100 mW/m². Therefore, the temperatures generated are a strong function of crater diameter.

Despite this wide range in temperature, simulations have shown that a significant proportion of the ejecta will not exceed 100 °C due to the existence of a ‘spall’ zone (Melosh 1984). This zone comprises much of the surface layer undergoing the impact and refers to a region in which the shock wave from the impactor is effectively cancelled via superposition with its reflected counterpart. These relatively low-temperature fragments would offer more favourable conditions to any lifeforms residing upon them. The following section will discuss specific forms of life that have shown great promise when tested against the panspermia hypothesis, from initial ejection (Section 5.1) through interplanetary travel (Section 5.2) to eventual atmospheric entry upon reaching the target planet (Section 5.3).

5. SURVIVAL OF MICRO-ORGANISMS

Over the past few decades, it has become possible to simulate the three stages of panspermia: (1) initial ejection from the impacted planet, (2) the subsequent journey through interplanetary space, and finally (3) impact with the target planet. Each step provides a new set of challenges to the survival of life. This section aims to supply a brief account of how certain micro-organisms have fared when placed in environments that are reminiscent of the panspermia process, and places these constraints into an extrasolar planetary context. A more comprehensive review of the near-Earth and Solar system contexts can be found in [Horneck et al. 2010].

5.1. Planetary ejection

A precondition for ejecta to be produced is the existence of impactors. Not every exoplanet host star contains compact asteroid belts (Martin & Livio 2013). However, evidence from white dwarf planetary systems indicate that between one quarter and one half of Milky Way white dwarfs host asteroid belts or Kuiper belt analogs (Koester et al. 2014) that are dynamically excited by a number of mechanisms involving planets (Veras 2016), the same fraction of main sequence stars thought to host planets throughout the Galaxy (Cassan et al. 2012). Further, the transfer of either biomolecules (pseudo-panspermia; Linkan & Loeh 2017a) or dead organisms (necropanspermia; Wesson 2016) imposes less stringent requirements than the transfer of living organisms.

In compact extrasolar systems, we might expect billions of rocks to be transferred between their planets, as it has been estimated that in the much more expansive solar system hundreds of millions of rocks could have already been ejected from the spall zones of Martian impacts and made their way to Earth (Mileikowsky et al. 2003). Such a healthy estimate has provided motivation for a number of investigators to conduct simulations which attempt to investigate the survivability of an impact large enough to eject material from Mars. One study, by Stöffler et al. (2007), applied pressures of 5 - 50 GPa to micron-thin layers of different microorganisms. This pressure range, applied via high explosives, is thought to be typical of Martian impact ejection (Artemieva & Ivanov 2004). Spores of Bacillus subtilis exhibited survival fractions (N/N₀ where N is the number of surviving cells and N₀ is the original number of viable cells) of 10⁻⁴ under a pressure of 42 GPa.

Bacterial spores are resilient casings that contain identical genetic information to their corresponding microorganism. Thought to be due to their low water content, the cores of bacterial spores have been found to exhibit notably low enzyme activity. This property is believed to contribute to the resilience of the spores, alongside the fact that the bacterial DNA is mixed with acid-soluble proteins, which aid in enzymatic reactivation (Setlow 1995). Besides the application of high explosives, the planetary ejection stage has also been simulated by firing projectiles at layers of spores, with similar survival rates to that outlined above (Burchell et al. 2004; Fajardo-Cavazos et al. 2009).

5.2. Journey through interplanetary space

Following successful ejection from the impacted planet, the life must then endure the trip to the target planet. Although not as violent as impact-driven ejection, this stage of interplanetary panspermia has been shown to be equally as deadly for a number of micro-organisms, owing to the deleterious conditions present in the space environment. The space between planets is vast and empty; vacuum pressures can drop as low as 10⁻¹⁴ Pa, inducing severe desiccation. Exposure to both stellar and Galactic cosmic radiation can also be highly damaging, with stellar UV posing the biggest threat. Temperature extremes during interplanetary transit have the potential to rival those of the ejection phase, depending on the orientation of the fragment’s orbit around its host star. Together, these form a lethal concoction for many microorganisms. Some, however, have exhibited impressive resilience against the harsh conditions of space.

The seven planets of TRAPPIST-1 orbit an ultra-cool M dwarf star. Due to the star’s relatively low temperature, its habitable zone lies very close in (at hundreds of an au). As such, it is likely to play host to a radiation environment that is much more damaging than the one surrounding the Earth. Investigations into the X-ray/EUV irradiation of the planets were undertaken by Wheatley et al. (2017). They deduced that TRAPPIST-1 has an X-ray luminosity similar to that of the quiet Sun; it has been hypothesised that such a high flux at these wavelengths could have stripped the planets of their atmospheres (Dong et al. 2017; Roettenbacher & Kand 2017), raising severe doubts regarding their habitability. The planetary atmospheres are also expected to alter frequently due to persistent flaring events (Vida et al. 2017), which can penetrate magnetospheres (Garraffo et al. 2017) and also affect the surface-based biospheres (Carone et al. 2018) in conjunc-
tion with geochemistry [Barr et al. 2017]. It would appear, therefore, that the TRAPPIST-1 planets are unlikely to support atmospheres that would enable the harbouring of life.

For more Solar system-like exoplanetary systems, we can consider the numerous exposure missions that have taken place in low Earth orbit (LEO), namely at altitudes less than 2000 km. It is important to note that the results of LEO experiments provide a mere estimate of the survivability of panspermia; such orbits are relatively close in proximity to the Earth, and therefore fail to accurately mirror the conditions expected during an interplanetary transit. For instance, the minimum vacuum pressure in LEO is approximately $10^{-6} - 10^{-7}$ Pa, several orders of magnitude higher than would be experienced for the majority of a planet-planet trip. Furthermore, the magnetic field of the Earth would shield the lifeform from much of the harmful cosmic radiation that would be plentiful in other regions of the Solar System. Nevertheless, LEO missions have contributed vastly to our understanding of the survival limits of copious micro-organisms, placing constraints on the plausibility of panspermia as a concept.

5.2.1. Long Duration Exposure Facility

Still holding the record for the longest exposure to LEO, NASAs Long Duration Exposure Facility (LDEF) subjected spores of B. subtilis to a combination of the space vacuum, solar UV and multiple components of Galactic cosmic radiation [Horneck et al. 1994]. In accordance with several other astrobiological experiments, solar UV was found to cause most damage due to its tendency to target and break DNA strands within the spores. Prospects for survival were greatly improved with adequate shielding from the UV in place; around 70% of B. subtilis spores were able to survive 6 years of exposure to the LEO vacuum when mixed with the sugar glucose.

5.2.2. EURECA

Similar conclusions were drawn from the results of the EURECA mission, which reported a complete loss of viability of Deinococcus radiodurans cells following 9 months of exposure to LEO [Dose et al. 1995]. Up to 12 DNA double strand breaks were observed per chromosome in samples exposed to the solar UV, although shielded cells also showed complete inactivation. These findings were surprising, as D. radiodurans is known to be incredibly resistant to desiccation and radiation; the bacterium can withstand radiation doses of 5000 Gray (around 1000 times a typically lethal dose for humans) with no loss of viability [Moseley et al. 1971].

A more recent study has found that D. radiodurans cells can survive many more dessication-induced DNA double strand breaks than those observed in the EURECA mission, due to the fact that the genome is able to reassemble before each successive cycle of cell division [Cox & Battista 2002]. As such, the survival rate observed following exposure to LEO conditions could have been higher than that inferred from the experimental results. Regardless, it is clear that the combination of stellar UV radiation and space vacuum form a deadly cocktail, survivable only for the most resilient of lifeforms known to inhabit our Earth. Indeed, vacuum-induced dehydration has been found to alter DNA photochemistry in such a way as to enhance the UV sensitivity of B. subtilis spores ten-fold in comparison to irradiation at atmospheric pressure [Horneck 1998; Nicholson 2000].

5.2.3. Biopan

Owing to these early findings, a general consensus has emerged that adequate shielding from the harmful environment of interplanetary space must be in place for micro-organisms, such as bacterial spores, lichens and tardigrades, to stand a chance of surviving panspermia.

Shielding — Mileikowsky et al. 2000 provide a thorough investigation of shielding for the case of Earth and Mars, and Cockell & Knowland 1999 provide a thorough summary of shielding mechanisms from UV radiation. The effects of shielding were explored as part of the series of experiments that took place using the Biopan facilities aboard various Foton satellites [Horneck et al. 2001]. A survival fraction of $10^{-6}$ was obtained when B. subtilis spores were exposed to the full LEO environment, whilst much higher fractions of 0.5 - 0.97 were determined for shielded samples. Clay shielding was found to be ineffective when placed in the form of a ‘shadowing’ layer; much more protection was received if the spores were mixed in with the clay, or ground meteorite powder. Importantly, the samples consisted of multilayers of spores; the outer layers would have encountered the full extent of LEO conditions, inactivating quickly and potentially forming a protective ‘crust’, offering added protection to the innermost layers of spores. It is also thought that endolithic micro-organisms, residing in microcracks present within rocks, likely exist in the form of biofilms embedded within a complex matrix of sugar molecules [Costerton et al. 1987]. This configuration would provide additional protection against the space vacuum. As such, lifeforms mixed within a layer of rock or clay are likely to receive much greater shielding from both the stellar UV and the space vacuum.

Lichens — During the Biopan 5 mission, thalli of the lichens Xanthoria elegans and Rhizocarpon geographicum were exposed to the space vacuum and selected wavebands of the solar UV for 14.6 days [Sancho et al. 2007]. A lichen comprises a stable symbiotic interaction between fungi and/or cyanobacteria. Lichens can be endolithic, growing between the grains inside rock, and are commonly found in mountainous regions. They have been found to survive complete water loss throughout periods of severe desiccation [Kramer et al. 2008] and withstand higher than average levels of UV radiation. Following the exposure to LEO, 83% of X. elegans cells were found to have intact membranes, whilst a similarly high survival rate of 71% was determined for R. geographicum.

Furthermore, full photosynthetic recovery was observed, even for samples exposed to over 99% of the solar light. The lichens contain certain pigments that provide screening from the UV, heightening protection during exposure, such as parietin phenolic acids [Solhaug et al. 2003]. Similar UV-screening properties were exhibited by cells of the halophilic cyanobacterium Synechococcus following two weeks of exposure to LEO as part of the Biopan 1 series of experiments [Mancinelli et al. 1998].
Interestingly, X. elegans has also been tested in simulations of the planetary ejection stage of panspermia. The lichen fared similarly to B. subtilis spores, with survival rates dropping by only four orders of magnitude upon the application of 50 GPa pressure (Horneck et al. 2008).

**Tardigrades** — Biopan 6, on the other hand, provided the first testing ground for tardigrades in space (Jönsson et al. 2008). Tardigrades have been identified as one of the most resilient animals on Earth, so are a natural choice for testing in LEO. They have been found to survive extreme temperatures and pressures for significant periods of time (Hengherr et al. 2009; Horikawa et al. 2009), and show incredible resistance to radiation, surviving doses of up to 5000 Gray (Hashimoto et al. 2010). A recent study by Sloan et al. (2017) deduced that tardigrades are likely to survive any mass extinction event with an astrophysical cause, such as a nearby supernova, gamma-ray burst or large asteroid impact.

In a similar way to bacterial spores, tardigrades can undergo a process known as cryptobiosis, whereby metabolic processes shut down in a reversible fashion during times of extreme stress. One particular form of cryptobiosis, known as anhydrobiosis, is of particular relevance to our discussion of survival in space. In this process, a tardigrade will contract and lose the vast majority of its water content, enabling cell stabilisers like trehalose to be formed and metabolism to, in the majority of its water content, enabling cell stabilisers like trehalose to be formed and metabolism to, in the most extreme cases, be temporarily halted altogether (Welnicz et al. 2011). Samples of the tardigrade species Milnesium tardigradum and Richtersius coronifer survived exposure to the LEO vacuum very well. Combined exposure to both the space vacuum and solar/Galactic radiation resulted in reduced, yet still finite, survival for both species tested. Tardigrades therefore have joined bacterial spores and lichens in the list of lifeforms that have survived exposure to the full LEO environment.

### 5.2.4. EXPOSE

The most recent results obtained from exposure missions in LEO are those of the European Space Agency’s EXPOSE facilities, mounted aboard various modules of the International Space Station. Conducted upon EXPOSE-E, the LIFE experiment subjected a variety of eukaryotic organisms to long-term exposure (1.5 years) for the first time (Onofri et al. 2012). Most notably, X. elegans once again achieved full photosynthetic recovery, provided the samples were shielded from UV irradiation. The AMINO experiment, which took place aboard the EXPOSE-R facility, exposed organic molecules to LEO both in their natural state and embedded in meteorite powder (Bertrand et al. 2013). Chosen for the key roles they play in the formation of macromolecules considered essential for life, the amino acids glycine, alanine and aspartic acid showed minimal deterioration following exposure, with 72 % of glycine remaining unaffected in unshielded form. Samples of the prokaryote Halorubrum chaoviarior, a halophilic archaeon, were exposed to LEO as part of the OSMO experiment (Mancinelli 2013). If shielded from the solar UV, the archaea exhibited 90 % survival rates.

### 5.3. Atmospheric entry

From the many exposure experiments that have taken place in LEO, it is clear that the deleterious conditions in space can have a devastating effect on many microorganisms. However, it is also clear that a number of lifeforms possess the necessary resilience to survive in such hazardous environments, especially when adequate shielding is in place. We now turn our attention to the final stage of material transfer: atmospheric entry upon reaching the target planet.

Because entry speeds range from 12 to 20 kms$^{-1}$ for typical asteroids, the overall process can occur in the space of a few tens of seconds (Nicholson 2009). Frictional heating over this rapid timescale leads to the formation of a fusion crust on the surface of the meteorite. This crust ensures that the heating fails to penetrate further than the first few millimetres of material, allowing the interior to maintain a relatively constant temperature throughout. Provided the target planet possesses an atmosphere, the eventual impact with the surface will occur at terminal velocity (50 m s$^{-1}$ for Earth), a far tamer value than what is involved in the planetary ejection phase.

Thus far, the best attempts to assess the ability of micro-organisms to survive meteoric entry have been those of the STONE missions, conducted upon the recovery module heat shields of the same Foton satellites used to host the Biopan 5 and 6 facilities (Parnell et al. 2008; Fouucher et al. 2010). The entry speed was measured to be 7.7 kms$^{-1}$, falling short of the expected speeds for asteroids provided above. Nevertheless, none of the micro-organisms tested showed any signs of viability following retrieval, most notably B. subtilis. For one of the samples, the fusion crust was found to be around 5 cm deep, possibly due to cracks in the surface of the shield. It would seem, therefore, that further experimentation is required to make any sort of conclusion regarding the survivability of the entry stage of panspermia.

### 6. CONCLUSIONS

The strong prospects for future discoveries of habitable multi-planet systems prompted us to analyze several aspects of panspermia and derive new results. Here, we have applied an impulse formalism from Jackson et al. (2014) to generate orbital constraints on life-bearing ejecta travelling between planets in the coplanar circular case (equations 1 and 3). Resulting analytic probability distributions depend only on the semimajor axes of the source and target planets (equations 11 and 12) and can be readily applied to compact multi-planet systems. We have also repackaged and consolidated physical relations that are associated with ejecta to fit within one framework (minimum radius and mass to escape atmosphere; equations 12–13; largest impact fragment: equations 13 and 14; minimum impactor size to liberate material: Section 4.3; speed and impactor radius to destroy source: equation 21). We finally included biological constraints from impact, interplanetary travel and atmospheric entry (Section 5). We hope that our results will represent useful tools to analyze future discoveries of compact multi-planet habitable systems.

### 7. ACKNOWLEDGMENTS

We thank both referees for particularly helpful and specific comments on the manuscript, resulting in an
APPENDIX

APPENDIX A: ORBIT TRANSFERS

In this appendix we provide formulae referenced in Section 2 and whatever additional prescriptions are needed to derive results. First, the orbit of the ejecta are related to the source planet through (Jackson et al. 2014)

\[
\frac{a}{a'} = 1 - \left(\frac{\Delta v}{v_k}\right)^2 - \frac{2}{\sqrt{1-e^2}} \left(\frac{\Delta v}{v_k}\right) \sin \theta \left[\sin \phi - e \sin \phi\right],
\]

(A1)

\[
e'' = 1 - (1-e^2) \left(\frac{H a}{a'}\right),
\]

(A2)

\[
\cos I' = \frac{1}{\sqrt{H}} \left[1 + \frac{\sqrt{1-e^2}}{1+e \cos f} \left(\frac{\Delta v}{v_k}\right) \sin \theta \sin \phi - f\right],
\]

(A3)

\[
\sin f' = \frac{\sqrt{H}}{e'} \left[\sin f + \sqrt{1-e^2} \left(\frac{\Delta v}{v_k}\right) \sin \cos \phi - f\right],
\]

(A4)

and

\[
\cos f' = \frac{1}{e'} [H (1 + e \cos f) - 1],
\]

(A5)

where

\[
H = 1 + 2 \frac{\sqrt{1-e^2}}{1+e \cos f} \left(\frac{\Delta v}{v_k}\right) \sin \theta \sin \phi + \frac{1-e^2}{(1+e \cos f)^2} \left(\frac{\Delta v}{v_k}\right)^2 \left[\cos^2 \theta + \sin^2 \theta \sin^2 \phi - f\right],
\]

(A6)

and where \(e'\) and \(f'\) refer to the eccentricity and true anomaly of the ejecta orbit. We also define the pericentre and apocentre of the ejecta orbit as

\[
q' = a' (1 - e'),
\]

(A7)

\[
Q' = a' (1 + e').
\]

(A8)

For coplanar and circular orbits, the pericenter \((q')\) and apocentre \((Q')\) of the debris’ orbit are constrained by

\[
q' \leq a \left[\frac{1 - \sqrt{X}}{1 - \left(\frac{\Delta v}{v_k}\right)^2 - 2 \left(\frac{\Delta v}{v_k}\right) \sin \phi}\right],
\]

(A9)
\[ Q' \geq a \left[ \frac{1 + \sqrt{\chi \left( \frac{\Delta v}{v_k} \right)}}{1 - \left( \frac{\Delta v}{v_k} \right)^2 - 2 \left( \frac{\Delta v}{v_k} \right) \sin \phi} \right], \]  

(A10)

where

\[ \chi = 1 + \sin \phi \left[ 2 \frac{\Delta v}{v_k} + \sin \phi \left( 3 + \left( \frac{\Delta v}{v_k} \right)^2 + 2 \sin \phi \frac{\Delta v}{v_k} \right) \right]. \]  

(A11)

The possible extrema of equation (1) are, for inward panspermia,

\[ (\phi)_{\text{ex}} = \{ \pm \frac{\pi}{2}, \pm \arccos \left[ \pm \frac{\sqrt{a^2 - q'^2}}{a} \right], \pm \arccos \left[ \pm \frac{\sqrt{2a^3 + 2a^2q' + 2aq'^2 + q'^3}}{a\sqrt{2a + q'}} \right] \}. \]  

(A12)

For outward panspermia, a similar set of extrema hold (just substitute \( q' \) for \( Q' \)) except for the middle ones, where the radicand would always be negative. The resulting minimum, real and nonzero values of \( \Delta v/v_k \) can then be converted into a minimum kick speed. For outward panspermia, the only real solutions are \( (\phi)_{\text{ex}} = -\frac{\pi}{2} \).

**Probability distributions**

In order to obtain the probability distribution functions for the debris to intersect the orbit of a planet, we first invert equations (1)-(3) as an explicit function of \( \sin \phi \) and write this expression in terms of known dimensionless ratios. Doing so yields

\[ (\sin \phi)_{\text{in}} = \frac{\alpha^2 - 1 \pm \sqrt{\alpha^2 (2 + \alpha^2 + \alpha (\gamma^2 - 3))}}{\gamma}, \]  

(A13)

\[ (\sin \phi)_{\text{out}} = \frac{1 - \beta^2 \pm \sqrt{2\beta^3 + 1 + \beta^2 (\gamma^2 - 3)}}{\gamma\beta^2}, \]  

(A14)

where

\[ \alpha \equiv \frac{q'}{a} < 1 \]  

(A15)

\[ \beta \equiv \frac{a}{Q'} < 1, \]  

(A16)

\[ \gamma \equiv \frac{\Delta v}{v_k}, \]  

(A17)

Equations (A13) and (A14) produce physical results only if the absolute value of the right-hand-sides are less than or equal to unity.

**Outward panspermia** — For outward panspermia – the simpler case – this condition is consistent with those found in equations (5-6) for only the lower sign in equation (A14). That equation therefore gives two values of \( (\sin \phi)_{\text{out}} \), denoted as \( \phi^1_{\text{out}}, \phi^2_{\text{out}} \), as long as \( \gamma \) is not sufficiently large or small. Assume over the interval [0, 2\( \pi \)] that \( \phi^1_{\text{out}} \leq \phi^2_{\text{out}} \). Consequently, for outward panspermia, we obtain

- **In the case of low kick speeds:** if

\[ \gamma < \sqrt{\frac{2}{1 + \beta}} - 1, \quad \text{then } P_{\text{out}} = 0. \]  

(A18)

- **In the case of high kick speeds:** if

\[ \gamma \geq \sqrt{\frac{2}{1 + \beta}} + 1, \quad \text{then } P_{\text{out}} = 1. \]  

(A19)
• In the case of medium kick speeds: if

\[
\sqrt{\frac{2}{1+\beta}} - 1 \leq \gamma < \sqrt{\frac{2}{1+\beta}} + 1, \quad \text{then}
\]

\[
P_{\text{out}} = \frac{1}{2\pi} \left\{ 2\pi \left\lfloor \frac{\phi_{\text{out}}}{\pi} \right\rfloor + (-1)^{\lfloor \phi_{\text{out}}/\pi \rfloor} \left( \phi_{\text{out}}^2 - \phi_1^{\text{out}} \right) \right\}.
\]

(A20)

This last equation represents visually the normalized horizontal distance between points on a curve from Fig. 3. The potential modulation by \(2\pi\) depends on whether or not this distance lies leftward or rightward of \(\phi = \pi\).

**Inward panspermia**— Now we perform a similar analysis for inward panspermia, which is more subtle. Consider again equation (A13). The condition that the absolute value of the right-hand-side of the equation is less than or equal to unity now does not exclude either sign in front of the radicand.

The equation with the upper sign is valid when \(\gamma \geq 1 - \sqrt{2\alpha/(1+\alpha)}\) and the equation with the lower sign is valid when \(\gamma \geq 1 + \sqrt{2\alpha/(1+\alpha)}\). The latter accounts for the “bump” seen around \(\phi = 270^\circ\) on Fig. 2. The former specifies points along the sloped portions of the other curves. The vertical lines on the figure correspond to extrema of \(\phi\).

Now let \(\phi_3^{\text{in}}\) and \(\phi_4^{\text{in}}\) be the solutions of equation (A14) with the lower sign, such that \(\phi_3^{\text{in}} \leq \phi_4^{\text{in}}\) on a \([0, 2\pi]\) interval. Consequently, we obtain

• In the case of low kick speeds: if

\[
\left[ \gamma < 1 + \sqrt{\frac{2\alpha}{1+\alpha}} \quad \text{and} \quad \alpha < \frac{1}{2} \left( \sqrt{2} - 1 \right) \right] \quad \text{or} \quad
\]

\[
\left[ \gamma < \frac{1 - \alpha}{2\alpha (1+\alpha)} \quad \text{and} \quad \alpha \geq \frac{1}{2} \left( \sqrt{2} - 1 \right) \right],
\]

then \(P_{\text{in}} = 0\). \hspace{1cm} (A21)

• In the case of medium kick speeds and wide separations, if

\[
1 + \sqrt{\frac{2\alpha}{1+\alpha}} \leq \gamma < \frac{1 - \alpha}{2\alpha (1+\alpha)} \quad \text{and} \quad \alpha < \frac{1}{2} \left( \sqrt{2} - 1 \right),
\]

then \(P_{\text{in}} = \frac{\phi_4^{\text{in}} - \phi_3^{\text{in}}}{2\pi}\). \hspace{1cm} (A22)

• In the case of medium kick speeds and close separations, if

\[
\frac{1 - \alpha}{2\alpha (1+\alpha)} \leq \gamma < 1 + \sqrt{\frac{2\alpha}{1+\alpha}} \quad \text{and} \quad \alpha \geq \frac{1}{2} \left( \sqrt{2} - 1 \right),
\]

then \(P_{\text{in}} = \frac{1}{2\pi} \left[ 2 \left( -\arccos \left( -\sqrt{1 - \alpha^2} \right) - \phi_1^{\text{in}} \right) \right]\). \hspace{1cm} (A23)

• In the case of high kick speeds: if

\[
\left[ \gamma \geq \frac{1 - \alpha}{2\alpha (1+\alpha)} \quad \text{and} \quad \alpha < \frac{1}{2} \left( \sqrt{2} - 1 \right) \right] \quad \text{or}
\]
\[ \gamma \geq 1 + \sqrt{\frac{2\alpha}{1 + \alpha}} \quad \text{and} \quad \alpha \geq \frac{1}{2} (\sqrt{2} - 1), \] then

\[ P_{in} = \frac{1}{2\pi} \left[ 2 \left( -\arccos \left( -\sqrt{1 - \alpha^2} \right) - \phi_1^{in} \right) + \phi_4^{in} - \phi_3^{in} \right]. \] (A24)

Equations (A21)-(A24) all represent normalized horizontal distances between a curve on Fig. 2. Because the curves feature multiple extrema, equations (A23)-(A24) in particular may not be obvious, but are derived from the same principle: if and where there are intersections between a curve and a horizontal line on the plot.

**APPENDIX B: A FIDUCIAL PACKED SYSTEM**

The motivation for the formalism presented in this paper is compact planetary systems that are within the habitable zones of their parent stars. Compact systems are said to be “packed”. The tendency for some systems to be packed was hinted at a decade ago (Barnes & Raymond 2004; Barnes et al. 2008; Raymond et al. 2009), but has now become commonplace thanks to the Kepler mission. In this appendix, we generate a compact planetary system which allows one to quickly compute characteristic values without relying on system-specific data.

In order for compact systems to be dynamically stable, the planets must be sufficiently separated from one another. Because no exact analytical prescription exists for three or more planets, many investigations have adopted empirical expressions. A commonly-used formulation is to express the separation of two planets through equations (B1-B2) of Chatterjee et al. (2008) as

\[ a_u = a_{u-1} + K \left[ \frac{M_{u-1} + M_u}{3M_{\text{star}}} \right]^{1/3} \left( \frac{a_{u-1} + a_u}{2} \right). \] (B1)

Here \( K \) is a constant and the planet subscripts are ordered in increasing distance from their star. The larger the value of \( K \), the longer the system will remain stable. However, as \( K \) increases, the scatter in the instability times becomes larger as well. For three-planet systems \( K \geq 5.5 \) generally ensures that planetary systems will remain stable for at least \( 10^9 \) yr.

If all of the planet masses are equal, then the recursive equation (B1) can be solved for arbitrary \( u \), yielding

\[ a_u = a_1 \left[ \frac{2 + K M}{2 - K M} \right]^{u-1}, \quad u \geq 1 \] (B2)

where

\[ M \equiv \left[ \frac{2M}{3M_{\text{star}}} \right]^{1/3} \] (B3)

Therefore, in an \( N \)-planet packed system with equal planet masses and circular orbits, the ratios \((a/Q')\) and \((a/q')\) are given just by a function of the constants \( K, M \) and the difference of the order numbers of the source and target planets. For example, suppose the source is planet \( j \) and the target, which is further away, is planet \( k \). Then

\[ \min (\Delta v)_{jk}^{\text{out}} = \sqrt{G (M_{\text{star}} + M)} a_1 \left[ \frac{2 + K M}{2 - K M} \right]^{1/3} \sqrt{2} \left( \frac{1 + \left[ \frac{2 + K M}{2 - K M} \right]^{j-k}}{2 - K M} \right)^{-1} - 1 \] (B4)

Consider, for example, a set of \( N = 6 \) Earth-mass planets orbiting a Solar-mass star. Then, \( M = 0.0126 \). Also assume that \( K = 7 \) provides sufficiently distant spacing for the planets to remain stable long enough to develop intelligent life. Consequently, \((2 + K M)/(2 - K M) = 1.092\). The escape speed from each planet is 11.19 km/s, and the circular speed of the innermost planet is \( 94.19 \text{ km/s} \times (a_1/0.10 \text{ au})^{-1/2} \). Note that with this last relation, all of the minimum thrusts computed may be scaled accordingly to the innermost semimajor axis. For \( a_1 = 0.10 \) au, the minimum kick speeds needed to reach the outermost planet from the other planets starting with the innermost are then \{43.9, 37.7, 31.3, 24.5, 16.6\} km/s.