Biological Homochirality and the Search for Extraterrestrial Biosignatures

Marcelo Gleiser

Received: 31 March 2022 / Accepted: 9 June 2022 / Published online: 15 August 2022
© The Author(s), under exclusive licence to Springer Nature B.V. 2022

Abstract
Most amino acids and sugar molecules occur in mirror, or chiral, images of each other, knowns as enantiomers. However, life on Earth is mostly homochiral: proteins contain almost exclusively L-amino acids, while only D-sugars appear in RNA and DNA. The mechanism behind this fundamental asymmetry of life remains unknown, despite much progress in the theoretical and experimental understanding of homochirality in the past decades. We review three potential mechanisms for the emergence of biological homochirality on primal Earth and explore their implications for astrobiology: the first, that biological homochirality is a stochastic process driven by local environmental fluctuations; the second, that it is driven by circularly-polarized ultraviolet radiation in star-forming regions; and the third, that it is driven by parity violation at the elementary particle level. We argue that each of these mechanisms leads to different observational consequences for the existence of enantiomeric excesses in our solar system and in exoplanets, pointing to the possibility that the search for life elsewhere will help elucidate the origins of homochirality on Earth.

Keywords Early planetary environments · Homochirality · Prebiotic chemistry · Origin of life

Introduction: A Bit of History
In 1815, the French physicist and chemist Jean-Baptiste Biot discovered that when light travelled through liquid solutions made out of a number of naturally occurring organic products, its polarization was affected. Pasteur was well aware of Biot’s studies Gleiser (2010). As he wrote in a set of lecture notes from 1860, “[Biot] quite definitely concluded that the action produced by the organic bodies was a molecular one, peculiar to their ultimate particles and depending on their individual constitution.” Pasteur (1848) The “action” Pasteur referred to was the ability of these natural organic compounds to rotate the polarization direction of light. With remarkable prescience, Biot had conjectured that such property was related to something going on at the molecular level. Pasteur put Biot’s conjecture...
into firm ground, showing that the optical properties of certain organic compounds—the way they interacted with light—resulted from the spatial structure of their individual molecules. Building upon Biot's research, Pasteur established that when linearly polarized light passed through a solution of tartaric acid synthesized in the lab, nothing happened: the synthetic solution was optically inactive. But when polarized light passed through a solution containing acid extracted from grapes, and thus from a living entity, its polarization direction changed.

Pasteur realized that since both substances had identical chemical properties, their molecules had the same types of atoms. What then could cause such puzzling asymmetric behavior? Could living and nonliving substances, even if apparently identical, have different properties? He examined the crystals from both substances under a microscope. He noted that whereas the lab-synthesized acid had two kinds of crystals, the acid from grapes had only one. With tremendous patience, he separated samples of both crystals using tweezers. Passing light through two solutions made with each of them, he demonstrated that the different crystals rotated the polarization plane of light in opposite directions: “I carefully separated the crystals which were [asymmetric] to the right from those [asymmetric] to the left, and examined their solutions separately in the polarizing apparatus. I then saw with no less surprise than pleasure that the crystals [asymmetric] to the right deviated the plane of polarization to the right, and that those [asymmetric] to the left deviated it to the left; and when I took an equal weight of each of the two kinds of crystals, the mixed solution was indifferent towards the light in consequence of the neutralization of the two equal and opposite individual deviations.” Pasteur (1848)

Pasteur's remarkable finding was that the naturally-occurring compound only appears in one of its two possible forms while the synthetic one appears in both. Was life selecting a specific molecular orientation? Continuing with his investigation, Pasteur showed that many organic compounds extracted from living organisms had the same biased optical properties. In one experiment, he added mold to a synthetic sample of tartaric acid. Initially, there was no optical activity, as expected. But as the mold grew, so did the optical activity of the sample. Furthermore, the increasing rotation was in the same direction of the naturally occurring acid. There was only one possible conclusion: life had a molecular bias. As Pasteur later wrote, “The Universe is dissymmetric and I am persuaded that life, as it is known to us, is a direct result of the asymmetry of the Universe or of its indirect consequences.”

Think of proteins as long chains of amino acids, pearl necklaces where each pearl is a molecular building block. Imagine that a left-handed (levorotatory) amino acid is a white pearl and a right-handed (dextrorotatory) amino acid a black pearl. Life has a clear (but not exclusive) preference for white pearl necklaces: the crucial molecules for life, proteins, are built from asymmetric backbones. The same is true for the sugar backbones of RNA and DNA. However, in this case the bias goes the opposite way: the sugars are dextrorotatory. It is hard to avoid the suspicion that this molecular bias is somehow related to the origin of life itself. Pasteur was the first to speculate as such: “Why even right or left substances at all? Why not simply non-asymmetric substances; substances of the order of inorganic nature? There are evidently causes for these curious manifestations of the play of molecular forces.... Is it not necessary and sufficient to admit that at the moment of the elaboration of the primary principles in the vegetable organism, an asymmetric force is present?” Pasteur (1848)

Pasteur had essentially posed the question we are still asking, whether homochirality has a specific biological function that, somehow, is related to some yet unknown fundamental causal mechanism. Although books Janoschek (1991); Wagnière (2007); Hochberg
Mechanisms for Chiral Bias

This section reviews three potential abiotic mechanisms for promoting chiral bias at the molecular level. Each will have a specific range of astronomical impact and thus be of relevance for astrobiological research and future searches for organic materials in our solar system and, via remote observation, in star-forming regions and exoplanets that display promising biosignatures. The author apologizes beforehand for the unevenness of the discussion, weighted disproportionately toward some of his work. However, with the added references and discussions, the interested reader can certainly pursue further details.

Punctuated Chirality: Planetary Bias

In reference Gleiser et al. (2008), Gleiser, Thorarinson, and Walker (GTW) proposed a mechanism dubbed “Punctuated Chirality” whereby the drive toward homochirality on Earth was the product of random environmentally-driven fluctuations that critically affected the prebiotic, molecular-forming mix of organic compounds, possibly many times over.
The starting point was Sandars’ polymerization model Sandars (2003), a generalization of Frank’s pioneering approach Frank (1953), featuring autocatalysis with enantiomeric cross-inhibition. Consider a left-handed polymer \( L_n \), made of \( n \) left-handed monomers, \( L_1 \). It may grow by adding another left-handed monomer with a rate \( k_s \), or be inhibited by adding a right-handed monomer \( D_1 \) with a rate \( k_I \). (Note that we denote D-compounds by the letter “D” as opposed to the notation set in Sandars’ work where such molecules were denoted as “R”.) The reaction network for \( n = 1, \ldots, N \), where \( N \) is the maximum polymer length in the system, can be written as:

\[
\begin{align*}
L_n + L_1 &\rightarrow_{2k_s} L_{n+1}, \\
L_n + D_1 &\rightarrow_{2k_I} L_n D_1, \\
L_1 + L_n D_1 &\rightarrow_{k_s} L_{n+1} D_1, \\
D_1 + L_n D_1 &\rightarrow_{k_I} D_1 L_n D_1,
\end{align*}
\]

supplemented by reactions for \( D \)-polymers by interchanging \( L \leftrightarrow D \), and by the production rate of monomers from the substrate:

\[
\begin{align*}
S \rightarrow_{k_C} L_1; \quad S \rightarrow_{k_C} D_1; \quad C_{L(D)} \text{ determine the enzymatic enhancement of } L(D)-\text{handed monomers, usually assumed to depend on the largest polymer in the reactor pool, } C_{L(D)} = L_N(D_N) \text{ Sandars (2003), or on a sum of all polymers Wattis and Coveney (2005). Soai’s group obtained the best-known illustration of this autocatalytic mechanism with enantiomeric cross-inhibition Soai et al. (1995), with dimers } (N = 2) \text{ as catalysts Blackmond (2004).}
\end{align*}
\]

A set of coupled, nonlinear ordinary differential equations for the various concentrations, \([L_1], [D_1], \ldots, [L_n], [D_n]\), describes the time evolution of the reaction network of Eq. (1), supplemented by the equation for the substrate, \( d[S]/dt = Q(Q_L + Q_D) \), where \( Q \) is the substrate’s production rate, and \( Q_L - Q_D = k_C f[S](C_L - C_D) \) gives the net chiral excess in monomer production. \( f \) is the enzymatic fidelity, usually set to unity to maximize chiral separation. GTW showed that starting as racemates, numerical solutions for polymerization reactions with \( N = 2, 5, \) and \( \infty \) evolve toward homochirality. This is also the case for \( N = 2 \) within the adiabatic approximation, where the rate of change for dimers and the substrate is assumed to be much slower than that of monomers, that is, when \( k_{S,L} << k_C \) Brandenburg and Multamäki (2004). In Gleiser and Walker (2008), a detailed study of the polymerization reaction network for various values of \( N \) has shown that the trends remain true when the effects of spatial dynamics are considered. Gleiser and Walker also concluded that although the adiabatic approximation predicts faster approach to steady-state conditions when compared with the full \( N = 2 \) model, it does produce the correct asymptotic values for the various concentrations.

Extending this network to include spatio-temporal diffusion—and thus departing from the well-mixed limit described by ODEs—starts by substituting \( d/dt \to \partial/\partial t - k \nabla^2 \), where \( k \) is the diffusion constant Brandenburg and Multamäki (2004). In this coarse-grained approach, the number of molecules per unit volume is large enough so that the concentrations vary smoothly in space and time. The spatiotemporal evolution of the network is obtained by solving the coupled system of nonlinear PDEs for arbitrary values of \( n \). Clearly, as \( n \) increases, solving and statistically analyzing the coupled system of equations in two and three spatial dimensions for various parameters becomes highly CPU intensive. Following Soai et al. (1995) where dimers were shown to be efficient catalysts (see also ref. Brandenburg and Multamäki (2004)), GTW focused on the
truncated system for $N = 2$ within the adiabatic approximation since, as with spatially-independent reaction networks, it has similar qualitative behavior to networks with longer ($N > 2$) polymer chains Gleiser and Walker (2008). The system then reduces to two coupled PDEs for the concentrations $|L_1|$ and $|D_1|$, being thus more amenable to a detailed statistical study while maintaining the key qualitative features of a larger reaction network. Typical results are shown in the top three panels of Fig. 1, where the two-phase system evolving from near-racemic conditions gets “stuck” due to the presence of both chiral phases.

The situation changes dramatically when noise is added to the system, as shown by GTW Gleiser et al. (2008). This was motivated by experimental demonstrations that stirring can bias chirality Kondepudi et al. (1990); Viedma (2005), and numerical studies for a similar $N = 2$ systems that explored how fluid turbulence can speed up chiral evolution Brandenburg and Multamäki (2004). Of course, the term “punctuated” makes reference to the “punctuated evolution” theory of Eldredge and Gould (1972), where random, intense phenomena of widespread environmental impact resets the evolutionary soup, so to speak, followed by longer periods of stasis. The role of the noise is to simulate random but substantial environmental perturbations that can affect the evolution of the biomolecular reaction network. In particular, the noise can, in principle, flip the direction of the chiral bias. This is achieved in the simplest possible

![Fig. 1](image-url) Evolution of 2d chiral domains. Red (+1 on the color bar) corresponds to the $L$-phase and blue (-1 on the color bar) corresponds to the $D$-phase. Time runs from left to right and top to bottom. Top left, the near-racemic initial conditions. Top mid and top right, evolution of the two percolating chiral domains separated by a thin domain wall. Bottom left, environmental effects break the stability of the domain wall network. Bottom right, subsequent surface-tension driven evolution leads to a enantiomerically-pure world Gleiser et al. (2008)
way via a generalized spatiotemporal Langevin equation Gleiser et al. (2006). The dynamical equations are written as:

\begin{equation}
\begin{align*}
\mathcal{L}_0^{-1}\left( \frac{\partial S}{\partial t} - kV^2S \right) &= 1 - S^2 + w(t, x), \\
\mathcal{L}_0^{-1}\left( \frac{\partial A}{\partial t} - kV^2A \right) &= S\mathcal{A} \left( \frac{2fS^2 + A^2}{S^2 + \mathcal{A}^2} - 1 \right) + w(t, x),
\end{align*}
\end{equation}

where \( \mathcal{L}_0 \equiv (2kS)^{1/2} \), and \( w(x, t) \) is a dimensionless Gaussian white noise with two-point correlation function \( \langle w(x', t')w(x, t) \rangle = a^2 \delta(t' - t)\delta(x' - x) \), where \( a^2 \) is a measure of the environmental influence’s strength. An Ising phase diagram can be constructed showing that \( \langle A \rangle \to 0 \) for \( a > a_c \); chiral symmetry is restored Gleiser et al. (2006). The value of \( a_c \) has been obtained numerically in two \( \left( a_c^2 = 1.15(k/l_0^2) \right) \) and three \( \left( a_c^2 = 0.65(k/l_0^2)^{1/2} \right) \) dimensions Gleiser et al. (2006). Dimensionless time, \( t_0 = l_0t \), and space, \( x_0 = x(l_0/k)^{1/2} \), variables were introduced. For diffusion in water \( k = 10^{-5}\text{m}^2\text{s}^{-1} \) and nominal values \( k_5 = 10^{-23}\text{cm}^3\text{s}^{-1} \) and \( Q = 10^{15}\text{cm}^{-3}\text{s}^{-1} \), we obtain \( l_0 = \sqrt{2} \times 10^{-5}\text{s}^{-1} \), simulating a 2d (3d) shallow (deep) pool with linear dimensions of \( l \sim 200 \) (50) cm. For the purpose of illustration, explicit results quoted below were computed using these values. Details of the numerical implementation can be obtained in Gleiser et al. (2008).

Results of a large statistical sample of 100 2d runs that led to initial domain coexistence, that is, \( d\langle A \rangle/dt \approx 0 \) show that near the critical region \( a^2 \geq 0.96a_c^2 \), all but the shortest events \( (t \leq 50l_0^{-1} \approx 1.5 \text{ months} \) for the nominal value of \( l_0 = \sqrt{2} \times 10^{-5}\text{s}^{-1} \) mentioned previously) lead to statistically significant chiral biasing. Results in 3d are qualitatively very similar, although due to heavy CPU demand we limited the analysis to 50 short runs. Essentially, large environmental disturbances modeled here as Gaussian noise with a certain amplitude and duration can not only drive a near-racemic system toward homochirality but also can reverse the chirality of homochiral solutions, effectively erasing any previous chiral signature.

As argued in GTW, the results suggest that, in the one extreme the early Earth may have played host to numerous abiogenetic events, only one of which ultimately led to the Last Universal Common Ancestor through the usual processes of Darwinian evolution. This is consistent with work indicating the widespread diversity and impact of extinction events, including the possibility of life emerging more than once Wilde et al. (2001). In the other extreme, one may consider, at the very least, that biological precursors certainly interacted with the primordial environment and may have had their chirality reset multiple times before homochiral life first evolved. In this case, our results show that separate domains of molecular assemblies with randomly set chirality may have reacted in different ways to environmental disturbances. A final, Earth-wide homochiral prebiotic chemistry would have been the result of multiple interactions between neighboring chiral domains under mechanisms described elsewhere Gleiser (2007); Gleiser and Walker (2008); Brandenburg and Multamäki (2004). If punctuated chirality prevails, it implies that homochirality in different planetary platforms conducive to life or, at least, to stereochemistry is a random process, and thus localized. This means that a statistically large sample of extraterrestrial stereochemistry would, on average, show no chiral bias. The Universe, taken as a whole, would be racemic.
Laboratory investigations have shown that enantiomeric excesses can be produced by asymmetric photolysis or synthesis Griesbeck and Meierhenrich (2002); Meierhenrich et al. (2005). On the other hand, active star-forming regions can generate circularly polarized light (CPL) Bailey (1998, 2001); Cataldo et al. (2005); Lucas et al. (2005). In this case, and if the process is efficient, the stereochemistry of all worlds (meaning planets and moons) within this region should have a clear prevalence of one chiral bias over the other. The discovery of an enantiomeric excess of chiral organic compounds in the Murchison meteorite has supported this view for our solar system (Pizzarello and Cronin 1998; Glavin and Dworkin 2005). Furthermore, examination of isotopic distributions (in $^{15}$N/$^{13}$N) and of α-branched amino acids of extraterrestrial origin has eliminated the possibility of contamination by Earth’s biosphere, with measurements showing an excess of L-alanine of 50% and of 30% for L-glutamic acid Engel and Macko (1997). A plausible explanation for such abiotic enantiomeric excess is that the star-forming region that originated the solar system was subjected to CPL, such as synchrotron radiation from a neutron star, although neutron stars don’t appear to be a significant source of CPL in the visible and UV range. If CPL-induced chiral bias is indeed a viable mechanism, the same chiral bias should be prevalent throughout the solar system but not necessarily throughout the galaxy. For example, finding an excess in D-chiral compounds in (Kminek et al. 2000; Glavin et al. 2020) or elsewhere in the solar system would contradict this scenario. (It would also contradict the possibility of a dominant chirality throughout the Universe, as we discuss in the next subsection.)

Chiral biasing through CPL depends on several unknowns such as the nature of the UV source, its distance from the target planet or moon, and the duration of effective radiative emission, making its viability harder to estimate Bailey (2001); Kondepudi and Nelson (1985). Furthermore, photolysis of amino acids requires UV radiation, which cannot be directly observed due to dust obscuration. Fukue et al. have investigated the range of CPL in the Orion nebula star forming region Fukue et al. (2010). Although they reported a high circular polarization region with considerable spatial extent ($\sim 0.4$ pc; for comparison, the distance between the Sun and Proxima Centauri is 1.3 pc) around the massive star-forming region known as the BN/KL nebula, other regions, including the linearly polarized Orion bar, show no significant circular polarization. Most of the low-mass young stars did not show detectable extended structure in either linear or circular polarization, in contrast to the BN/KL nebula. So, although CPL is a potential mechanism for generating an early bias toward enantiomeric excess, the only way to confirm its viability is by probing several worlds within the same original star-forming region. Certainly, finding meteorites with the same enantiomeric excess as on Earth strengthens this hypothesis Pizzarello and Cronin (1998); Glavin and Dworkin (2005). However, the situation remains ambivalent. For example, Glavin and Dworkin didn’t detect L-isovaline excess for the pristine Antarctic CR2 meteorites Elephant Moraine 92042 and Queen Alexandra Range 99177, whereas they did detect large levorotatory enantiomeric excesses in the CM meteorite Murchison and the CI meteorite Orgueil Glavin and Dworkin (2005). They suggest that this excess was produced by the amplification of a small initial excess by an aqueous alteration phase.

Another obstacle to CPL-induced enantiomeric excess is its resilience against environmental disturbances as discussed above. A clear signature of a CPL-induced enantiomeric excess is that it must be correlated with a planet’s formation era, as the meteoritic
evidence suggests. If punctuated chirality is viable, it becomes difficult to explain the long-term resilience of a CPL-induced excess during, in the case of our planet, a window of a few hundred-million years, when the terrestrial environmental was certainly prone to large-scale perturbations due to heavy bombardment and active volcanism. Convincing evidence that CPL is the preferred mechanism for generating an enantiomeric excess would require a systematic search in most of our solar system. If, indeed, the same handedness prevails across our solar system, then CPL becomes a viable possibility. However, such evidence shouldn’t be considered as the prevalent mechanism operating across the galaxy, unless parity-violation in the weak interactions can indeed be ruled out. We examine this mechanism next.

Chirality from Parity Violating Interactions

The discovery of parity violation in the weak interactions, and the subsequent formulation of electroweak theory in the Standard Model of particle physics suggests the possibility that parity-violating forces could affect quantum chemical calculations, including perturbative computations of parity violating potentials which, in turn, could play a role in biasing chirality at the molecular level Yamagata (1966); Salam (1991); Kondepudi and Nelson (1985). This would be an impressive connection between dynamics at the smallest level of elementary particle physics and the mechanisms that drive the homochirality of living systems, being, of course, of great aesthetic appeal. If this were the mechanism for biasing biological homochirality, prebiotic stereochemistry across the Universe would have to be the same, unifying the physics of the very small with cosmic stereochemistry and, possibly, life on Earth with life everywhere else in the Universe (if any).

Although initial results were small to the point of being negligible, more detailed calculations led to parity-violating energy differences of \(\sim 10^{-11} \text{ J mol}^{-1}\) (approximately 100 aeV) in enantiomers of chiral molecules Bakasov et al. (1998); Quack et al. (2008). Clearly, even with this increase in the effect, a very efficient amplification mechanism is needed in order to affect molecular interactions of interest for biochemistry.

Kondepudi and Nelson (1985) constructed an ODE model where a small enantiomeric excess can be amplified by parity violation and is also subjected to CPL fluctuations. Unfortunately, their initially optimistic results were put into question when a more detailed spatiotemporal analysis was developed Gleiser (2007). This work used a similar parameterization as ref. Kondepudi and Nelson (1985) to express the bias due to parity violation in the weak nuclear interactions, which has been estimated to be \(g = E_f/k_B T \sim 10^{-17-18}\) at room temperature (Salam 1991; Lazzeretti et al. 1999; Bakasov et al. 1998). In ref. (Gleiser 2007) it was shown that using the well-mixed limit when discussing the dynamics greatly enhances the time scales where the chiral excess amplification can be effective. Indeed, including spatial dependence of the concentrations implies that the dynamics of symmetry breaking can take two possible paths, depending on the relative volume occupied by each phase: in the first case, if both phases are above percolation threshold, the system evolves as the domain walls separating the two phases vie for dominance. This domain-wall dynamics is clearly quite different from that of simple ODEs for the concentrations of enantiomers, and models the early Earth environment more realistically. Indeed, setting as a realistic time-scale for chirality to be defined as 100 million years implies that the parity-violating biasing must satisfy \(g \geq 7 \times 10^{-6}\) (100My/\(t_g\))\(^{1/2}\) (Gleiser 2007), unfortunately more than ten orders of magnitude larger than the quantum chemistry computations. For \(g \leq 10^{-6}\) it would take longer than the age of...
the Universe before biasing becomes active in moving the walls toward a single phase. An estimate of the velocity with which the walls propagate shows that for a wall to convert a distance of 1km in 100My, \( g \geq 4 \times 10^{-6} \). So, it not only takes too long for the chiral bias \( g \) to take over the dynamics of the domain wall system, but, once it does take over, the walls move too slowly to sweep an effective area in a realistic time scale \( \sim 100\text{My} \).

In the second case, a two-phase system with one phase in a metastable state (the one unfavored by the chiral bias), and the walls moving slowly, we can examine how the dynamics evolves through bubble nucleation typical of first-order phase transitions (Gleiser 2007). Unfortunately, the situation is worse in this case. If we impose the same time-scale for bubble nucleation as above, \( \tau_{\text{nuc}} \leq 100\text{My} \), a “shallow” cylindrical pool with volume \( \sim \pi \times 10^8\text{m}^3 \), gives \( \tau_{\text{nuc}} \simeq 4 \times 10^{-18} \exp[89/g^2a^2]\text{y} \). Choosing a fairly large value \( a^2 \simeq 0.5 \) for the typical noise amplitude that induces thermal fluctuations (cf. Fig. 2), we obtain \( g \geq 1.74 \), an unrealistically large value. One possibility left open is whether external perturbations could accelerate the decay of the metastable state. These could involve a variety of external influences, from cataclysmic events to meteoritic impact, perhaps already with some enantiomeric excess, thus combining punctuated chirality with phase transition dynamics.

Although we cannot discard future advances in the simulation of phase dynamics or other potential amplification mechanisms such as those just mentioned, it seems unlikely at this point that parity violation played a role in biasing the biological homochirality observed on Earth. A strong test of this hypothesis would be to have a statistically representative sample of stereochemistry over many stellar systems. If the chiral bias toward a specific enantiomeric excess for all these samples is the same as on Earth, then we may infer that indeed parity violation has found a way to influence chirality across the Universe. The only caveats, of course, are, one, whether punctuated chirality is effective at a planetary scale and, two, whether unknown alien biological processes can also be effective in reversing chirality. If so, either or both of these processes would erase any memory of a prior uniform prebiotic chiral excess with varying levels of effectiveness across different planetary platforms, and we would be back to chirality as being contingent to a planet’s specific geophysical history or biological processes.

---

**Fig. 2** Punctuated Chirality
Gleiser et al. (2008). Impact of environmental effects of varying duration and fixed magnitude \( (a^2/a_0^2 = 0.96) \) on the evolution of prebiotic chirality in 2d. Short events (last from left), which have little to no effect, should be contrasted with longer ones, which can drive the chirality towards purity and/or reverse its trend. (See, e.g. the green line.)
Concluding Remarks: Observing Biological Chirality in the Universe

As mentioned above, clarifying the abiotic mechanism or mechanisms responsible for biasing a specific enantiomeric excess on Earth or elsewhere will rely on data collected from other planetary platforms, in this solar system and others Avnir (2021). The fundamental premise, inspired by what we know from life on Earth, is that even if the initial chiral biasing mechanism is abiotic, biological processes will incorporate this excess and take it to the next level. In terrestrial proteins, enzymatic function is dependent on the folding of amino acid chains into highly ordered structures, which is not possible without a chiral excess. Furthermore, since there are no known abiotic sources of enantiomeric excesses of D-amino acids or L-sugars on Earth, finding such excesses in another solar system planet or moon would be a strong indication of extraterrestrial biological processes Glavin et al. (2020). Such a find would also support the hypothesis of punctuated chirality operating to generate distinct enantiomeric excesses at individual planetary platforms.

Several in situ and sample return missions are now in progress or under development. NASA’s Curiosity rover Creamer et al. (2016) and the Philae lander of the Rosetta spacecraft Meierhenrich et al. (2013) were both equipped with gas-chromatographic coupled to mass spectrometry instruments capable of detecting enantiomeric excesses. Although at present in situ measurements of complex organics with enantioselective chromatography (coupled to a variety of detectors) are challenging to current spaceflight instrumentation, returning samples to Earth may be the best avenue to methodically and reliably search for enantiomeric excesses in extraterrestrial soil. Recent successful sample collection by the Perseverance rover opens the possibility that, when returned and analyzed on Earth by the end of the current decade, will allow the identification of chiral organic compounds on the Martian soil. The ESA, China, Japan, and Russia also have plans for sample collection and return.

Another avenue for the identification of extraterrestrial chiral compounds is through remote sensing, such as the recent discovery of propylene oxide in the Sagittarius B2 star-forming region using a radio telescope McGuire et al. (2016). This was the first chiral molecule discovered in outer space. Finding a chiral molecule in a star-forming region raises the prospect that chiral materials are available during the formation of proto-planetary disks, thus possibly being incorporated in nascent planets and diverse orbiting bodies such as asteroids and comets. This may well have been the case for our solar system.

Radio astronomy is the primary method for studying the complex molecular content of interstellar clouds, including spectral features corresponding to fine-structure transitions of atoms or pure rotational transitions of polar molecules. In principle, such methods at high-precision and full polarization can even detect enantiomeric excesses, although that was not the case for ref. McGuire et al. (2016). With this resource, we should hope for exciting results in the coming years where target star-forming regions could, in principle, reveal chiral organics with noticeable enantiomeric excesses.

Looking further ahead, a statically significant sample of diverse star-forming regions with chiral organics will be invaluable in the determination of the mechanism(s) driving the enantiomeric excess. If they all show the same skewness, a universal causation mechanism acting across the galaxy (and possibly the Universe) such as parity violation would be strongly favored. Otherwise, CPL acting on different star-forming clouds would be favored, generating the same chiral excesses for each parent star region but different skewness from region to region. Either way, we would have strong observational evidence that the molecules that jump-started the abiotic processes that led to biological homochirality on
Earth are operational across the galaxy. If such initial excess gets amplified or reversed due to environmental effects remain an open question. In this case, a large sample that tends to a neutral enantiomeric excess would support punctuated chirality or other unknown mechanisms with similar planetary-wide impact. Finally, looking much further ahead, and inspired by the recent successful launch of the James Webb Space Telescope, we could imagine a scenario where exoplanets with promising biosignatures have their transit spectra examined at high resolution for signs of chiral biomolecules in their atmosphere. If the same compounds found on Earth are identified, we would have strong evidence for biotic activity in the exoplanetary atmosphere, pointing toward universal biochemical properties. Whether such compounds would have similar or opposite chirality to their terrestrial counterparts would reveal the predominant abiotic mechanisms that drive homochirality here and elsewhere in the cosmos.

Acknowledgements The author wishes to thank the editors of the present volume for the invitation to contribute

Declarations

Conflict of interest The author declares no conflict of interest

References

Avnir D (2021) Critical review of chirality indicators of extraterrestrial life. New Astron Rev 92:101596
Bailey J (1998) Circular polarization in star-formation regions: implications for biomolecular homochirality. Science 281:672–674
Bailey J (2001) Astronomical sources of circularly polarized light and the origin of homochirality. Orig Life Evol Biosph 31:167–183
Bakasov A, Ha T-K, Quack M (1998) Ab initio calculation of molecular energies including parity violating interactions J Chem Phys 109:7263–7285
Blackmond DG (2019) The origin of biological homochirality. cold spring harbor perspectives in biology 11(3):032540
Blackmond DG (2004) Asymmetric autocatalysis and its implications for the origin of homochirality. PNAS 101:5732–5736
Brandenburg A, Multamäki T (2004) How long can left and right handed life forms coexist?. Int J Astrobiol 3:209–219
Cataldo F, Brucato JR, Kehey Y (2005) Chirality in prebiotic molecules and the phenomenon of photo- and radioracemization. J Phys Conf Series 6:139–148
Cline D (2000) The physical origin of homochirality in life (aip conference proceedings) aip conference proceedings 329
Creamer JS, Mora MF, Willis PA (2016) Enhanced resolution of chiral amino acids with capillary electrophoresis for biosignature detection in extraterrestrial samples. Anal Chem 89(2):1329-1337
Eldredge N, Gould SJ (1972) in Models in Paleobiology (ed. Schopf, T. J. M.) Ch. 5 (Freeman Cooper, San Francisco)
Engel MH, Macko SA (1997) Isotopic evidence for extraterrestrial non-racemic amino acids in the Murchison meteorite. Nature 389:265–268
Finefield JM et al (2012) Enantiomeric natural products: occurrence and biogenesis. Angew Chem Int Ed 51:4802–4836
Frank FC (1953) On spontaneous asymmetric catalysis. Biochim Biophys Acta 11:459–463
Fukue T et al (2010) Extended high circular polarization in the orion massive star forming region: implications for the origin of homochirality in the solar system. Orig Life Evol Biosph 40:335–346
Glavin DP, Dworkin JP (2005) Enrichment of the amino acid L-isovaline by aqueous alteration on CI and CM meteorite parent bodies. Proc Nat Acad Sci 106(14):5487–5492
Glavin DP et al (2020) The search for chiral asymmetry as a potential biosignature in our solar system. Chem Rev 120(11):4660–4689
Gleiser M (2007) Asymmetric spatiotemporal evolution of prebiotic homochirality. Orig Life Evol Biosph 37:235–251
Gleiser M (2010) A tear at the edge of creation: a radical new vision for life in an imperfect universe Free Press, New York, NY
Gleiser M, Thorarinson J (2006) Prebiotic homochirality as a critical phenomenon. Orig Life Evol Biosph 36:501–505
Gleiser M, Thorarinson J, Walker SI (2008) Punctuated Chirality. Orig Life Evol Biosph 38:499–508
Gleiser M, Walker SI (2008) An extended model for the evolution of prebiotic homochirality: a bottom-up approach to the origin of life. Orig Life Evol Biosph 38:293–315
Griesbeck AG, Meierhenrich UJ (2002) Asymmetric photochemistry and photochirogenesis. Angew Chem Int Ed 41:3147–3154
Hochberg D ed (2021) Asymmetry in biological homochirality Mdpi AG
Janoschek R (1991) Chirality from the Weak Bosons to the α-helix Springer Berlin Heildeberg, 1991.
Kminek G et al (2000) F. MOD: An organic detector for the future robotic exploration of Mars. Planet Space Sci 48:1087–1091
Kondepudi DK, Nelson GW (1985) Weak neutral currents and the origin of biomolecular chirality. Nature 314:438–441
Kondepudi DK, Kaufman RJ, Singh N (1990) Chiral symmetry breaking in sodium chloride crystallization. Science 250:975–976
Lazzeretti P, Zanasi R, Faglioni F (1999) Energetic stabilization of d-camphor via weak neutral currents. Phys Rev E 60:871–874
Lucas PW et al (2005) UV circular polarization in star formation regions: the origin of homochirality?. Orig Life Evol Biosph 35:29–60
McGuire BA et al (2016) Discovery of the interstellar chiral molecule propylene oxide CH3CHCH2O. Science 352(6292):1449–1452
Martin W, Russell MJ (2007) On the origin of biochemistry at an alkaline hydrothermal vent. Phil. Trans. R. Soc. B 362:1887–1925
Meierhenrich UJ, Muñoz C, Schutte WA, Thiemann WH-P, Barbier B, Brack A (2005) Precursors of biological cofactors from ultraviolet irradiation of circumstellar/interstellar ice analogs. Chem Eur J 11:4895–4900
Meierhenrich UJ et al (2013) Evaluating the robustness of the enantioselective stationary phases on the Rosetta mission against space vacuum vaporization. Adv Spac Res 52(12):2080–2084
Pasteur L. (1848) Recherches sur les relations qui peuvent exister entre la forme cristalline et la composition chimique, et le sens de la polarization rotatoire. Ann Chim Phys 24:442–459
Pizzarello S, Cronin JR (1998) Alanine enantiomers in the Murchison meteorite. Nature 394:236
Quack M, Stohner J, Willeke M (2008) High-resolution spectroscopic studies and theory of parity violation in chiral molecules. Ann Rev Phys Chem 59:741–769
Salam A (1991) The role of chirality in the origin of life. J Mol Evol 33:105–113
Sandars PGH (2003) A Toy model for the generation of homochirality during polymerization. Orig Life Evol Biosph 33:575–587
Soai K, Shibata T, Choji K, Morioka H (1995) Asymmetric autocatalysis and amplification of enantiometric excess of a chiral molecule. Nature 378:767–768
Viedma C (2005) Chiral symmetry breaking during crystallization: complete chiral purity induced by nonlinear autocatalysis and recycling. Phys Rev Lett 94:065504
Wagnière GH (2007) On chirality and the universal asymmetry : reflections on image and mirror image. VHCA with Wiley-VCH, Zürich
Wattis JA, Coveney PV (2005) Symmetry-breaking in chiral polymerization. Orig Life Evol Biosph 35:243–273
Wilde SA, Valley JW, Peck WH, Graham CM (2001) Evidence from detrital zircons for the existence of continental crust and oceans on the Earth 4.4 Gyr ago. Nature 409:175-178
Yamagata Y (1966) A hypothesis for the asymmetric appearance of biomolecules on earth J Theoret Biol 11:495–498

Springer