Enantiomeric excess by magnetic circular dichroism in Archaean atmosphere

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Evolution of homochirality requires an initial enantiomeric excess (EE) between right and left-handed biomolecules. We show that magnetic circular dichroism (MCD) of sun’s ultraviolet C light by oxygen in Archaean earth’s anoxic atmosphere followed by chirally selective damage of biomolecules due to circular dichroism (CD) can generate EE of correct handedness. Our calculation of EE uses published data for CD of biomolecules and accepted magnitude for Archaean earth’s magnetic field. Independent of atmospheric oxygen concentration calculated EE has the same sign for all pyrimidine nucleosides which is opposite to that for amino-acids. Purine nucleosides have smaller EE values with opposite sign to pyrimidines but are less susceptible to UV damage. Homochirality is explained by origin of prebiotic life in one hemisphere of earth and its evolution to EE ~ ± 1 before reversal of terrestrial magnetic field. Chirality of biomolecules is decided by the direction of magnetic field where prebiotic life originated on Archaean earth.

While homochirality is pervasive in molecules which form the building blocks of life1, results of Urey-Miller experiment2 form the basis of hypothesis that life originated on earth with a racemic mixture of prebiotic molecules. It is believed that evolution of homochirality rests on: (i) a mechanism to create initial enantiomeric excess (EE) between right (R) and left (L)-handed biomolecules and (ii) amplification of EE over time to generate 100% homochirality13. Two possible mechanisms have been proposed to explain the initial EE which evolved to 100% homochirality over a period of time. One of these has an extra-terrestrial source and rests on the observed small EE of L-amino acids found on chondritic meteorites4. However, it is possible that some of the evidence on these meteorites may be due to contamination5 by the L-amino acids of our biosphere. Second mechanism is stochastic and the observed homochirality is explained as a “by-chance” phenomenon6,7. Neither of these two mechanisms can explain why all five nucleosides are R-handed and all twenty standard amino acids (except achiral glycine) have the opposite L-handedness. Presence of terrestrial magnetic field can in principle break the chiral symmetry of biomolecules. In an effect labeled “magnetochiral dichroism”, unpolarized light propagating in the direction of a magnetic field can produce an EE in some chemical reactions involving metal and organic compounds8. However, its role in creating an EE for biomolecules like nucleic and amino acids remains to be demonstrated.

We show that initial EE of correct handedness can be generated by MCD of ultraviolet C (UVC) light by paramagnetic oxygen molecules in the atmosphere of Archaean earth followed by chirally selective photo damage due to CD in biomolecules. Dextro (right-handed) ribose and deoxyribose sugar is primarily responsible for the chirality of nucleic acids. These sugars have strong absorption/CD bands only for vacuum UV wavelengths of less than 190 nm9,10. As shown in this report, UV flux reaching Archean earth was limited to wavelengths above 200 nm making isolated sugar molecules immune to UV photolysis. Nucleosides are the smallest chiral building blocks of nucleic acids with strong circular dichroism for UV wavelengths in the 200–300 nm range and are analyzed together with amino-acids for EE. The model implies that enantiomeric enrichment by asymmetric photolysis with UV light happened after bigger molecules like nucleic acid and amino acids remains to be demonstrated.

Our analysis utilizes published data for CD of L-amino-acids and R-nucleosides and is based on the accepted understanding of paleomagnetism11 and Archaean atmosphere – 4.000 Mya. Evidence from Archaean rocks related to mass independent fractionation of sulphur (MIF-S) isotopes suggests an anoxic atmosphere12 with partial pressure of oxygen, p(O2) < 10^-2 PAL (present atmospheric level) although there are reports of loss of MIF-S and possible oxic atmosphere even 3.8 billion years ago13. While MIF-S effect still needs to be understood14 to make quantitative conclusions, based on photochemical modeling, maximum pressure of atmospheric
oxygen\textsuperscript{12} that would support MIF-S is $p(O_2) \sim 10^{-2}$ PAL. As shown in this report the sign of EE of nucleosides and amino-acids is independent of oxygen concentration but depends significantly on the pressure of carbon-dioxide in Archaean atmosphere.

**Results**

**Transmission of UVC in Archean atmosphere.** For analysis in this report, carbon dioxide and oxygen are the two most important components of Archaean atmosphere when prebiotic life arose around 4 billion years ago. While CO\textsubscript{2} is primarily responsible for attenuating far UV sunlight in wavelength range of 200–300 nm\textsuperscript{15}, O\textsubscript{2} is the only atmospheric component which is paramagnetic and capable of producing MCD of this UV light. Concentration of O\textsubscript{2} in atmosphere will determine the extent of MCD of UV light and the differential circular polarization (CP) intensity reaching earth's surface. Accepted models\textsuperscript{16} for earth's atmosphere around 4,000 Mya require high CO\textsubscript{2} pressure (0.1–10 bar) to compensate for lower solar luminosity of young Sun. Figure 1 is a plot of calculated UVC flux reaching early Archaean earth's surface ($p(CO_2) = 0.25, 1, 2$ bar) and shows that wavelengths below 200 nm were strongly attenuated. In plotting these curves, accepted values are used for CO\textsubscript{2} absorption cross-sections, density-altitude profile and the spectral-intensity of sunlight reaching earth's upper atmosphere 4,000 Mya\textsuperscript{15}. For wavelengths in 200–300 nm, molecular oxygen absorbs weakly by three forbidden transitions\textsuperscript{17,18}, giving rise to the Herzberg continuum which originates in paramagnetic $X^3\Sigma^−_g$ ground state and is important for atmospheric physics. As seen in Fig. 1, total absorption cross section for these transitions in oxygen ($X^3\Sigma^−_g \rightarrow A'^3\Sigma^+_u$, $X^3\Sigma^−_g \rightarrow c^1\Sigma^−_u$ and $X^3\Sigma^−_g \rightarrow A^'3\Delta_u$) rapidly decreases from 200 nm to 300 nm\textsuperscript{17–19}. In laboratory, MCD of O\textsubscript{2} has been observed\textsuperscript{20} within the Herzberg continuum by a matrix-isolation technique at low temperatures and high magnetic field. Attenuation by CO\textsubscript{2} and absorption cross-section for oxygen in Fig. 1 serves to explain why the effect of MCD by Archaean atmospheric oxygen is unimportant outside the spectral region of 200–300 nm.

**Magnetic circular dichroism of UVC in Archean atmosphere.** The dipolar geomagnetic field is as old\textsuperscript{11} as the Earth itself with the dipole axis nearly but not exactly aligned with axis of earth’s rotation. Paleomagnetism\textsuperscript{11,21} evidence suggests that the magnitude of magnetic field has not changed significantly since 4,000 Mya. The magnetic poles have wandered by as much as 15–20° but for the purpose of this analysis we will assume that the axis of dipole coincides with the axis of earth’s rotation (Fig. 2). We also assume the magnetic field intensity at the equator of 0.3 Gauss which is the present average value\textsuperscript{22} for this latitude; and the tilt of earth’s axis towards the ecliptic plane as $\theta_T = 23.4°$ (current value). For MCD, the component ($B_z$) of geomagnetic field parallel to the direction of sunlight reaching earth at zenith (midday) varies with the latitude as well as the time of the year and is derived with reference to Fig. 2a. Close to the surface of Earth (radius R), the radial ($B_r$) component of the dipolar magnetic field and the component parallel ($B_\parallel$) to earth’s surface is given by\textsuperscript{22},

$$B_r = -2B_0 \sin(\theta_L) \quad \text{and} \quad B_\parallel = B_0 \cos(\theta_L)$$

where $B_0$ is the magnetic field at the equator and $\theta_L$ is the latitude of the position on earth. Average value of $B_0$ on the equator is 0.3 Gauss.

For the geometry shown in Fig. 2a (northern solstice) when the North Pole is most tilted towards the sun, the component of earth’s magnetic field in the direction of sunlight at zenith (midday) is:

$$B_S = -B_r \cos(\theta_L - \theta_T) + B_\parallel \sin(\theta_L - \theta_T)$$

![Figure 1](image-url) Logarithmic plots of spectral flux of UVC light reaching earth’s surface for atmospheric CO\textsubscript{2} pressure of 0.25, 1 and 2 bar and of absorption cross section of molecular oxygen by the forbidden transitions of the Herzberg bands. The figure shows that the spectral region outside of 200–300 nm is unimportant for magnetic circular dichroism by Archean atmospheric oxygen.
where $\theta_T = 23.4^\circ$ is the angle of tilt of the earth’s axis towards the ecliptic plane. Similarly, for southern solstice when the North Pole is most tilted away from the sun, the component of earth’s magnetic field in the direction of sunlight at zenith (midday) is:

$$B_S = -B_I \cos(\theta_T + \theta_L) + B_I \sin(\theta_T + \theta_L)$$  \hspace{1cm} (3)

Both $B_I$ and $B_I$ vary\(^{22}\) with radial distance ($r$) as $(R/r)^3$ which shows that for distances of up to 100 km above earth’s surface $B_S$ varies by less than 3%. Atmosphere of this thickness accounts for 99% of all gaseous oxygen around earth\(^{22}\).

Magnetic field $B_S$ is shown in Fig. 2(a) for calculating $B_S$, the component of field in the direction of sunlight, (b). Variation of $B_S$ (component of geomagnetic field intensity in the direction of sunlight at zenith) with latitude for the two extremes of solstices. Dipolar magnetic field intensity of 0.3 Gauss at equator is assumed. Dark gray regions of latitude are relatively important for MCD where $B_S$ does not change sign throughout the year and $|B_S| > 0.25$ Gauss. For light gray regions $B_S$ changes sign in the course of a year and so are not important for MCD.

**Figure 2.** (a) Model of terrestrial magnetic dipole for calculating $B_S$, the component of field in the direction of sunlight. (b) Variation of $B_S$ (component of geomagnetic field intensity in the direction of sunlight at zenith) with latitude for the two extremes of solstices. Dipolar magnetic field intensity of 0.3 Gauss at equator is assumed. Dark gray regions of latitude are relatively important for MCD where $B_S$ does not change sign throughout the year and $|B_S| > 0.25$ Gauss. For light gray regions $B_S$ changes sign in the course of a year and so are not important for MCD.

The significance of MCD-related transitions\(^{20}\) in molecular oxygen which results in net circular polarization of UVC light is explained with Fig. 3. Of the three forbidden transitions, $X^3\Sigma_g^- \rightarrow A^3\Sigma_u^+$ is the strongest, accounting for most (86%) of the absorption\(^{18}\) in Herzberg continuum. Zeeman splitting of states by the small ($\sim 0.25$ Gauss) geomagnetic field and possible transitions with left circularly polarized (LCP, $\sigma^+$) and right circularly polarized (RCP, $\sigma^-$) light (propagating in the direction of magnetic field) are shown schematically. In the ground state $X^3\Sigma_g^-$, Boltzmann distribution will result in a small excess of population in $M_S = -1$ over $M_S = +1$. As seen in this figure, for $X^3\Sigma_g^- \rightarrow A^3\Sigma_u^+$ and $X^3\Sigma_g^- \rightarrow c^1\Sigma_u^-$ transitions, $M_S = -1$ in the ground state can only absorb $\sigma^+$ light while $M_S = +1$ can only absorb $\sigma^-$ light. Population difference between $M_S = -1$ and $M_S = +1$ results in slightly greater absorption of $\sigma^+$ light over $\sigma^-$. This is not possible with $X^3\Sigma_g^- \rightarrow A^3\Delta_u$ transition (Fig. 3). Net
(albeit very small) circular polarization of light reaching earth's surface is a result of $^1\Sigma \rightarrow \Sigma$ transitions in molecular oxygen, i.e. UV light reaching earth will be net $\sigma^-$ when propagating in the direction of earth's magnetic field and net $\sigma^+$ when propagating against the direction of magnetic field.

Differential (net) circular polarization (CP) intensity due to MCD by atmospheric O$_2$ is derived as follows: If $I_+ (\lambda)$ and $I_- (\lambda)$ is the spectral intensity (W.m$^{-2}$.nm$^{-1}$) of LCP ($\sigma^-$) and RCP ($\sigma^+$) circularly polarized solar UV radiation respectively, reaching the earth's surface after passing through the atmosphere,

$$I_+ (\lambda) = \frac{I_0 (\lambda)}{2} e^{-\alpha_+}$$ and $$I_- (\lambda) = \frac{I_0 (\lambda)}{2} e^{-\alpha_-} \quad (4)$$

$I_0 (\lambda)$ is the initial (above atmosphere) unpolarized intensity (equal components of RCP and LCP). Coefficients $\alpha_+$ and $\alpha_-$ describe bulk of the atmospheric absorption by CO$_2$ and relatively very weak differential absorption of circularly polarized light (MCD) by atmospheric O$_2$.

$$\alpha_+ (\lambda) = \sigma (\text{CO}_2, \lambda) \cdot \int N(\text{CO}_2, L) \cdot dL + \sigma_+ (\text{O}_2, \lambda) \cdot \int N_+ (\text{O}_2, L) \cdot dL \quad (5a)$$

Likewise,

$$\alpha_- (\lambda) = \sigma (\text{CO}_2, \lambda) \cdot \int N(\text{CO}_2, L) \cdot dL + \sigma_- (\text{O}_2, \lambda) \cdot \int N_- (\text{O}_2, L) \cdot dL \quad (5b)$$

The integration is over altitude ($L$) between 0 and 100 km of atmosphere which adequately includes absorption by more than 99% of CO$_2$ and O$_2$.

$\sigma (\text{CO}_2, \lambda)$: Absorption cross-section for CO$_2$ at wavelength $\lambda$

$N(\text{CO}_2, L)$: Number density of CO$_2$ at altitude L

$\sigma_+ (\text{O}_2, \lambda)$: absorption cross-section for O$_2$ at wavelength $\lambda$ (LCP light)

$\sigma_- (\text{O}_2, \lambda)$: absorption cross-section for O$_2$ at wavelength $\lambda$ (RCP light)

$N_+ (\text{O}_2, L)$: Number density of O$_2$ in $M_3 = -1$ at altitude L

$N_- (\text{O}_2, L)$: Number density of O$_2$ in $M_3 = +1$ at altitude L

For small magnetic field, $H \approx 0.25$ Gauss, the magnetic sublevels of O$_2$ are degenerate and we assume, $\sigma_- (\text{O}_2, \lambda) = \sigma_+ (\text{O}_2, \lambda) \equiv \sigma (\text{O}_2, \lambda)$

Boltzmann distribution gives,

$$\left( \frac{N_+ (\text{O}_2, L)}{N_- (\text{O}_2, L)} \right) = e^{\Delta E (H)/kT} \quad (6)$$

Here, $\Delta E (H) = \mu_B g \cdot \Delta M_3 H$ is the energy difference between $M_3 = -1$ and $M_3 = 1$ Zeeman levels for magnetic field $H; \mu_B$ is the Bohr magneton, $g$ is the Lande factor, $\Delta M_3 = 2$ for the two relevant sublevels of the ground state X$^3\Sigma_g^-$, and $k$ is the Boltzmann constant. We assume, $H \equiv B_0 = 0.25$ Gauss, Lande $g$ factor$^{25} \approx 2$ and $T = 300$ K giving, $\Delta E (H)/kT = 2.24 \times 10^{-2}$.

Due to three nearly degenerate magnetic sublevels of the ground state of O$_2$,

$$N_+ (\text{O}_2, L) = \frac{N (\text{O}_2, L)}{3} \quad \text{and} \quad N_- (\text{O}_2, L) = \frac{N (\text{O}_2, L)}{3} \cdot e^{-\Delta E (H)/kT} \quad (7)$$

$N (\text{O}_2, L)$ is the total number density of O$_2$ at altitude $L$. Spectral intensities $I_+ (\lambda)$ and $I_- (\lambda)$ are calculated from above equations, published data for absorption cross-sections $\sigma (\text{CO}_2, \lambda)$ and $\sigma (\text{O}_2, \lambda)$, and the density-altitude profile$^{15}$ for given surface pressure of CO$_2$. We assume complete mixing of CO$_2$ and O$_2$, i.e., $N (\text{O}_2, L)/N (\text{CO}_2, L)$ is independent of altitude $L$. Differential circular polarization (CP) intensity $\Delta I (\lambda) \equiv (I_+ (\lambda) - I_- (\lambda))$
is calculated from Eqns 4–7 and shown in Fig. 4 for a surface O₂ pressure p(O₂) = 0.002 bar and CO₂ pressure of 0.5 bar. Similar plots can be generated for any other surface O₂ and CO₂ pressure. As seen in this figure, ΔI(λ) is proportional to B_S / T.

**Enantiomeric excess for biomolecules.** Calculation of EE is made for biomolecules in aqueous medium as is the case with most theories for origin of prebiotic life in marine/freshwater environment. The products of Miller Urey type reactions with atmospheric gases collected over the large surface area of Archean oceans. Concentration of amino acids in Archean oceans is estimated to vary from 4 × 10⁻³ M to 10⁻⁷ M. Concentration of five nucleobases in one Miller-Urey experiment was measured to be between 1–100 ppm. While these numbers are small, there are several proposed mechanisms for amplifying the concentration of prebiotic molecules on ocean shorelines and drying freshwater ponds. Additionally, enantiomeric excess depends on the ratio and not absolute values of enantiomeric concentrations. As shown in a recent report, around 4000 Mya the pH of seawater was in the 6.5–7.0 range and not too different from freshwater. Data used in this report is for solutions of pH around 7. While asymmetric photolysis and enantiomeric enrichment of amino acids by circularly polarized light in liquid varies with the acidic pH in the range of 2–6, there is no effect of pH in the 6.5–7.0 range or higher.

Chiral molecules like nucleic acid monomers and amino acids show circular dichroism in the 200–300 nm region of UV light. Starting with a racemic population of these prebiotic building blocks of life, exposure to net circularly polarized UVC light will result in an enantiomeric excess that will be a seed for evolution to homochirality. Such an explanation has been proposed for the observed enantiomeric excess of L-amino acids in chondritic meteorites and assumes that the UV damage/inactivation rates D_R, D_L of right-/left-handed biomolecules are proportional to light absorption rates. This is also seen from the observed similarity of absorption spectrum and action spectrum for direct photo-damage of nucleic acids by UVC radiation. Using published CD parameter Δε(λ), extinction coefficient ε(λ), and calculated differential CP intensity ΔI(λ) = (I(+)(λ) − I(−)(λ)), the differential UV damage rate ΔD/D ≡ (D_R − D_L) / (D_R + D_L) for nucleosides and amino acids is derived below.

As shown here, enantiomeric excess is related to ΔD/D.

Cross sections for absorption of UV by biomolecules (amino-acids/nucleosides) are listed below.

σ_R⁺(λ): Cross-section for absorption of σ⁺ light by right-handed (RH) molecule
σ_R⁻(λ): Cross-section for absorption of σ⁻ light by right-handed (RH) molecule
σ_L⁺(λ): Cross-section for absorption of σ⁺ light by left-handed (LH) molecule
σ_L⁻(λ): Cross-section for absorption of σ⁻ light by left-handed (LH) molecule

For the two kinds of chiral molecules (RH/LH) and two circular light polarizations (σ⁺ and σ⁻), following rate equations are valid. LCP (σ⁺) energy absorbed per sec by one RH molecule is

$$I_R^+ = \int \sigma_R^+ (\lambda) \cdot d\lambda$$

Similarly, absorption rates for other three combinations,

$$I_R^- = \int \sigma_R^- (\lambda) \cdot d\lambda$$

$$I_L^+ = \int \sigma_L^+ (\lambda) \cdot d\lambda$$

$$I_L^- = \int \sigma_L^- (\lambda) \cdot d\lambda$$

Figure 4. Differential circular polarization intensity for p(O₂) = 0.002 bar and p(CO₂) = 0.5 bar. (a) ΔE/kT = 2.24 × 10⁻⁷, (b) ΔE/kT = 4.48 × 10⁻⁷.
\[ \Gamma^+_L = \int I_L(\lambda) \cdot \sigma^+_L(\lambda) \cdot d\lambda \]  
\[ \Gamma^-_L = \int I_L(\lambda) \cdot \sigma^-_L(\lambda) \cdot d\lambda \]  

(8c)

(8d)

Since chiral molecules can absorb both \( \sigma^+ \) and \( \sigma^- \) components of UV light, total absorption rates for RH and LH enantiomers are:

\[ \Gamma^+_R = \Gamma^+_R + \Gamma^-_R \]  
\[ \Gamma^-_L = \Gamma^+_L + \Gamma^-_L \]  

(9a)

(9b)

From chiral symmetry, \( \sigma^+_R(\lambda) = \sigma^-_L(\lambda) \) and \( \sigma^-_R(\lambda) = \sigma^+_L(\lambda) \). As before, differential circular polarization (CP) intensity due to MCD by atmospheric \( O_2 \) is \( \Delta I(\lambda) \equiv I_R(\lambda) - I_L(\lambda) \). Define, \( \Delta \sigma(\lambda) \equiv \sigma^+_L(\lambda) - \sigma^-_L(\lambda) = \sigma^+_R(\lambda) - \sigma^-_R(\lambda) \). It follows from above rate Eqs 8 and 9,

\[ \Gamma^-_R - \Gamma^+_L = \int \Delta I(\lambda) \cdot \Delta \sigma(\lambda) \cdot d\lambda \]  

(10a)

Similarly,

\[ \Gamma^+_R + \Gamma^-_L = \int I_R(\lambda) \cdot \sigma(\lambda) \cdot d\lambda \]  

(10b)

Here, \( I_L(\lambda) \approx I_R(\lambda) = I(\lambda)/2 \) and \( \sigma(\lambda) = \sigma^+_L(\lambda) + \sigma^-_L(\lambda) = \sigma^+_R(\lambda) + \sigma^-_R(\lambda) \). Molar circular dichroism is defined as \( \Delta \varepsilon(\lambda) \equiv \varepsilon^+ - \varepsilon^- \), where \( \varepsilon^+ \) and \( \varepsilon^- \) are the molar extinction coefficients (\( \text{mol}^{-1} \cdot \text{cm}^{-1} \)) for \( \sigma^+ \) and \( \sigma^- \) light respectively. Absorption cross-section \( \sigma(\lambda) \) is proportional\(^{20} \) to \( \varepsilon(\lambda) \). Thus,

\[ \frac{\Gamma^-_R - \Gamma^+_L}{\Gamma^+_R + \Gamma^-_L} = \frac{\int \Delta I(\lambda) \cdot \Delta \varepsilon(\lambda) \cdot d\lambda}{\int I_R(\lambda) \cdot \varepsilon(\lambda) \cdot d\lambda} = \frac{-\int \Delta I(\lambda) \cdot \Delta \varepsilon_R(\lambda) \cdot d\lambda}{\int I_R(\lambda) \cdot \varepsilon(\lambda) \cdot d\lambda} \]  

(11a)

Here, \( \Delta \varepsilon_R = \varepsilon^+_L - \varepsilon^-_L \) for L-handed molecule and \( \Delta \varepsilon_R = \varepsilon^+_R - \varepsilon^-_R \) for R-handed molecule. Since \( \Delta \varepsilon(\lambda) \) is typically less than 1% of \( \varepsilon(\lambda) \), in the denominator \( \varepsilon(\lambda) \approx \varepsilon^+_L(\lambda) \approx \varepsilon^+_R(\lambda) \). It is commonly assumed that the UV destruction/damage rate \( (D) \) for nucleic acids\(^{46,47} \) is proportional to the light energy absorption rate \( (\Gamma) \). Thus,

\[ \frac{\Delta D}{D} \equiv \frac{D_R - D_L}{D_R + D_L} = \frac{\int \Delta I(\lambda) \cdot \Delta \varepsilon_L(\lambda) \cdot d\lambda}{\int I_R(\lambda) \cdot \varepsilon(\lambda) \cdot d\lambda} = \frac{-\int \Delta I(\lambda) \cdot \Delta \varepsilon_R(\lambda) \cdot d\lambda}{\int I_R(\lambda) \cdot \varepsilon(\lambda) \cdot d\lambda} \]  

(11b)

Relative efficiency for photolysis has been studied only for a few amino acids and for some isolated wavelength. As an example, photolysis of phenylalanine at 206 nm and 254 nm does not show any significant effect of wavelength\(^{48} \). Further, CD of amino acids is significant only over a band-width of 20–30 nm\(^{34} \). Above equation to calculate \( \Delta D/D \) is used for nucleosides and amino acids.

In eqn. 11, \( \Delta \varepsilon_L(\lambda), \Delta \varepsilon_R(\lambda) \) is the molar CD parameter for L/R-handed enantiomer. Enantiomeric excess \( (N_R - N_L)/(N_R + N_L) \) is the relative excess of right-handed molecular concentration \( (N_R) \) over left-handed \( (N_L) \) molecules. It is related to \( \Delta D/D \) and the sign of EE is opposite to that of \( \Delta D/D \) (if \( D_R > D_L, N_R < N_L \)). In a simple model described below, for racemic production and UV damage of biomolecules enantiomeric excess, EE is \( -\Delta D/D \).

**Model for racemic production and UV damage of biomolecules.** Correlation between EE and the parameter \( (\Delta D/D) \) is explained with the help of a simple phenomenological model that describes the racemic production of prebiotic life-molecules and their destruction by Archaean UVC radiation. For number density \( N_R \) and \( N_L \) of right and left handed molecules,

\[ \frac{dN_R}{dt} = -D_R N_R + P \quad \text{and} \quad \frac{dN_L}{dt} = -D_L N_L + P \]  

(12)

The terms \( -D_R N_R - D_L N_L \) describe destruction by UV light with rate constants \( D_R \) and \( D_L \) which could be different due to differential circular polarization intensity, \( \Delta I(\lambda) \equiv I_R(\lambda) - I_L(\lambda) \). The racemic production rate \( P \) is same for both enantiomers. For time, \( t \gg D_R^{-1}, D_L^{-1} \), the steady-state distribution \( (dN_R/dt \approx 0) \) results in an enantiomeric excess,

\[ EE \equiv \frac{N_R - N_L}{N_R + N_L} = -\left( \frac{D_R - D_L}{D_R + D_L} \right) \equiv -\frac{\Delta D}{D} \]  

(13a)

For time, \( t \ll D_R^{-1}, D_L^{-1} \),

\[ EE \equiv \frac{N_R - N_L}{N_R + N_L} = -(D_R - D_L) \cdot \frac{t}{4} \equiv -\frac{\Delta D}{D} \left( \frac{Dt}{4} \right) \]  

(13b)
population density is racemic (EE ≈ 0, Dpyrimidine ≈ 45 and the corresponding damage time for pyrimidine nucleosides in the Archaean UV flux will be several hours. As seen from Equation 13a, b, for the estimated Archean UV flux, EE for pyrimidines will saturate to −ΔD/D in a time of few minutes but will take several hours for purines to do so. Compared to pyrimidines, the Archean purine population density is racemic (EE ≈ 0).

Relatively fewer studies have measured wavelength specific UV photolysis rate constants for amino acids. Photolysis of amino acids for wavelengths shorter than 200 nm was investigated for space environment outside of terrestrial atmosphere where UV light at these wavelengths is relatively abundant. Photolysis has also been investigated for wavelengths longer than 200 nm and these are the only ones relevant to our investigation due to absence of vacuum UV light on Archean earth. In one investigation 14, a comparison was made of the relative effectiveness of amino-acid photolysis at wavelengths of 147 nm, 206 nm and 254 nm. The results show that photolysis effectiveness at these three wavelengths is comparable. For UV flux of 2.66 W/m² at a wavelength of 254 nm, the measured half-life of phenylalanine was 130 min and for UV flux of 8.34 W/m² at a wavelength of 206 nm, it was 38 min. Without any wavelength specific damage action spectra for amino acids for 200–300 nm, it is difficult to compare UV photolysis rates for amino acids. However, the photolysis rate is likely to be higher for wavelengths longer than 200 nm, which is plausible due to the higher absorption of amino acids in this wavelength range. In general, amino acids are known to be more UV sensitive than nucleic acids.

Table 1. Summary of Circular Dichroism (CD) bands for L-amino-acids and R-nucleosides. Calculated values of EE for different pressures of atmospheric CO₂ and O₂ is given in Table 2 for nucleosides and five randomly selected amino-acids for a scenario where the intensity of σ⁺-light reaching earth's surface is more than σ⁻-intensity (ΔI(λ) ≡ I⁺(λ) − I⁻(λ) > 0).

Table 2. Calculated Enantiomeric Excess (EE) for atmospheric CO₂ pressure between 0.25–2 bar and an O₂ pressure of 0.002 bar. EE values are for I⁺(λ) > I⁻(λ), i.e. ΔI(λ) > 0 and when UVC light reaching earth's surface is preferentially σ⁻ (RCP).

Equation 11b is used to calculate EE (−ΔD/D) using published values of CD parameter Δε(λ) and extinction coefficient ε(λ) for R-nucleosides and L-amino-acids. A summary of reported CD bands for these biomolecules is given in Table 1.
is not possible to correlate these numbers with the Archaen UVC flux shown in Fig. 1. However, in the absence of any dramatic variation of photolysis rates with UVC wavelength, amino-acid half-life of a few hours is expected.

The UV absorption coefficient of water varies from 7 m\(^{-1}\) for a wavelength of 200 nm to 0.7 m\(^{-1}\) for 300 nm and has a value of 1.7 m\(^{-1}\) for the central wavelength of 250 nm. Thus UV light in this wavelength range easily penetrates around 0.5 meters of water. The products of Urey Miller type reactions will continue interacting with UV light till the molecules have sunk to depths greater than ~0.5 meters by diffusion. Diffusion coefficient (D) for molecules of the size of nucleosides and amino acids in water has a value of \(1.5 \times 10^{-5}\) cm\(^2\)/s. Time to diffuse down by a distance (L) of 0.5 m is \(t \approx L^2/4D\) several months. This can be compared to a time of several minutes to a few hours required for photodamage of biomolecules in the Archaean UV light flux, as explained above. Thus biomolecules in water get adequately illuminated for the photodamage processes described in this report with near-full UV flux from sun.

As seen from Table 1, \(\Delta \varepsilon_{\lambda}\) is positive for CD of L-amino-acids in 200–300 nm wavelength region. From equation 11b, \(\Delta D/D\) is positive for \(\Delta I(\lambda) > 0\). Thus EE is negative (\(N_l > N_R\)) for amino-acids and the sign of EE is independent of CO\(_2\) pressure (Table 2). Unlike amino-acids, the CD spectra of nucleosides show both positive and negative bands for \(\Delta \varepsilon_{\lambda}\) (Table 1). Increasing atmospheric CO\(_2\) shifts the UV pass-band (Fig. 1) and the differential circular polarization (Fig. 4) to longer wavelengths, resulting in a change in sign of \(\Delta D/D\) and (EE) for pyrimidine nucleosides. As seen in Table 2, for pyrimidine nucleosides EE is positive (\(N_R > N_L\)) above a threshold CO\(_2\) pressure of around 1 bar. In these calculations, the pressure of atmospheric O\(_2\) is kept at 0.002 bar (0.01 PAL). At low pressures of O\(_2\), calculated values of EE is approximately proportional to O\(_2\) pressure. This is shown (Table 2) for CO\(_2\) pressure of 1.25 bar and lower O\(_2\) pressure of 0.0002 bar. While magnitude of EE values reduce by a factor of 10, the sign of EE remains unchanged.

Interestingly, the magnitude of EE for both purine nucleosides is smaller and the sign is opposite to that for pyrimidines. However, as explained above (Eqn. 13), compared to pyrimidines EE will take much longer time to reach the saturation value of \(\Delta D/D\) and the population of purines is relatively racemic. Further, heterochiral nucleic acids involving pyrimidine and purine building blocks are significantly more unstable thermally as compared to homochiral counterparts. It is conceivable that smaller EE values for purines and significant thermal stability of homochiral nucleic acids together with \(D_{\text{purine}} \approx 0.01(D_{\text{pyrimidine}})\) was a determining factor (not the sign of their enantiomeric excess) in the evolution of chirality of purines. Values of \(\Delta D/D\) for aromatic amino-acids are lower by a factor of ~100 due to larger extinction coefficient \(\varepsilon(\lambda)\) and relatively small circular dichroism \(\Delta \varepsilon(\lambda)\). In all of these calculations we assume an ambient temperature (T) of 300 K and a magnetic field (B\(_s\)) of 0.25 Gauss. Varying conditions of atmospheric temperature and magnetic field can be accounted by the observation that differential circular polarization intensity \(\Delta I(\lambda)\) and EE are proportional to \(B_s/T\) (Fig. 4).

**Conclusions**

We have shown that magnetic circular dichroism of UVC by sparse oxygen in the anoxic Archaean atmosphere results in net circularly polarized light reaching earth's surface. This results in chirally selective damage of prebiotic molecules by circular dichroism and creates an EE which evolved to homochirality of R-nucleosides and L-amino-acids. Irrespective of the partial pressure of oxygen, the sign of enantiomeric excess of pyrimidine nucleosides and the amino-acids is predicted correctly when UVC light reaching earth has net \(\sigma^-\) circular polarization \((\Delta I(\lambda) \equiv I(\lambda) - I(-\lambda) > 0)\) and \(P(\text{CO}_2) > 1\) bar. This is consistent with the requirement of large CO\(_2\) pressure for greenhouse effect to balance lower solar luminosity 4,000 Mya. It is also an indication that initial enantiomeric excess was generated by circularly polarized UV light and not by a stochastic event.

Since MCD depends on the direction of magnetic field with respect to the propagation of light, net \(\sigma^-\) light in one hemisphere of earth would also result in net \(\sigma^+\) light in the other hemisphere. Homochirality in life-molecules can be explained by assuming (i) prebiotic life originated either in the northern or the southern hemisphere of earth which had higher UVC flux of \(\sigma^-\) over \(\sigma^+\) (south magnetic hemisphere), (ii) evolution of homochirality was complete (EE = ±1) before the earth's magnetic field reversed. Recorded length of polarity intervals between reversals varies between 0.1 and several Million years and (iii) no significant dispersal of prebiotic life molecules happened between hemispheres before the transition to homochirality was complete. For the terrestrial magnetic field in Fig. 2, the most important regions for effective MCD in atmospheric oxygen lie between latitudes of 24° and 58° in both hemispheres and any EE will be nullified by large-scale dispersal of enantiomers between these regions from south to north magnetic hemisphere.

**Methods**

**Calculation of spectral flux \(I(\lambda) (W.m^{-2}.nm^{-1})\) of UVC light reaching earth’s surface.**

\(I(\lambda) = \int I(\lambda) d\lambda \quad (W.m^{-2}.nm^{-1})\),

Equations 4 and 5 together with published data\(^1\) for the following is used:

1a. Spectral flux \(I_0(\lambda)\) (200–300 nm) from model young sun (~4,000 Mya) reaching earth (before entering the atmosphere)

1b. Absorption cross-sections (200–300 nm) for \(\text{CO}_2\)

1c. Variation of density with altitude (up to 120 km) for \(\text{CO}_2\) and \(\text{O}_2\). Altitude up 100 km accounts for 99% of these gases.

1d. Absorption cross-sections (200–300 nm) for \(\text{O}_2\) is from refs\(^17\-19\).

**Calculation of enantiomeric excess \((-\Delta D/D)\) from equation 11b.**

Published data (200–300 nm) for the following is used:

2a. Extinction coefficient \(\varepsilon\) (mol\(^{-1}\).cm\(^{-1}\)) for the four amino acids (Ala, Ser, Lys, Pro) is significant only for 200–230 nm. For aromatic amino acid (Phe), \(\varepsilon\) is significant between 200–270 nm.
2b. Circular dichroism (milli.deg) for amino acids is converted to \( \Delta\varepsilon \) (mol\(^{-1}\) cm\(^{-1}\)) and is significant only for 200–240 nm. Due to a small range of wavelengths for amino-acids, published data for \( \Delta\varepsilon \) and \( \varepsilon \) values is taken in steps of 1 nm.

Following equation (11b) is used to evaluate EE (\( -\Delta D/D \)) for amino-acids

\[
EE = -\frac{\Delta D}{D} = -\frac{\int I_0(\lambda) \cdot \Delta\varepsilon_0(\lambda) \cdot d\lambda}{\int I_0(\lambda) \cdot \varepsilon(\lambda) \cdot d\lambda} = -\frac{\sum_{\lambda=200}^{300} \Delta I(\lambda) \cdot \Delta\varepsilon(\lambda)}{\sum_{\lambda=200}^{300} I(\lambda) \cdot \varepsilon(\lambda)}
\]

Summation is done in steps of 1 nm.

2c. Extinction coefficient (mol\(^{-1}\) cm\(^{-1}\)) and circular dichroism (mol\(^{-1}\) cm\(^{-1}\)) for the five nucleosides

Due to a larger range of wavelengths for nucleosides, published data for \( \Delta\varepsilon \) and \( \varepsilon \) values is taken in steps of 2 nm.

Following equation (11b) is used to evaluate EE (\( -\Delta D/D \)) for five nucleosides

\[
EE = -\frac{\Delta D}{D} = -\frac{\int I_0(\lambda) \cdot \Delta\varepsilon_R(\lambda) \cdot d\lambda}{\int I_0(\lambda) \cdot \varepsilon(\lambda) \cdot d\lambda} = -\frac{\sum_{\lambda=200}^{300} \Delta I(\lambda) \cdot \Delta\varepsilon_R(\lambda)}{\sum_{\lambda=200}^{300} I(\lambda) \cdot \varepsilon_R(\lambda)}
\]

Summation is done in steps of 2 nm. As seen from Figs 1 and 4, there is very little UV light intensity for wavelengths shorter than 220 nm and there is insignificant contribution to EE from this wavelength region. Likewise, differential circular polarization intensity (Fig. 4) is minimal for wavelength longer than 290 nm and again there is insignificant contribution from this long-wavelength region.

**Data availability.** The author declares that the main data supporting the findings of this study are available within this article. Additional data used in this study is available publicly in published journal articles. Appropriate references are provided (Methods section) for the sources of this data.

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**Author Contributions**
A.S. conceived this study and wrote the paper.

**Additional Information**

**Competing Interests:** The authors declare that they have no competing interests.

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