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temperatures of the D["] layer, MgSiO₃ transforms from perovskite into a layered CaIrO₃-type post-perovskite phase. The elastic properties of the post-perovskite phase and its stability field explain several observed puzzling properties of the D["] layer: its seismic anisotropy², the strongly undulating shear-wave discontinuity at its top³⁻⁶ and possibly the anticorrelation between shear and bulk sound velocities^{7,8}.

If MgSiO₃ perovskite is stable throughout the lower mantle, it should be the most abundant mineral in our planet. While some researchers⁹ have suggested its decomposition into the oxides at lower mantle conditions, most workers^{10–13} have found that perovskite is more stable than the oxides. To our knowledge, the possibility that MgSiO₃ could be stable in a completely new structure within the lower mantle has not been considered.

The shear-wave discontinuity at the top of the D" layer, suggested in ref. 3, has a strong topography. This discontinuity has commonly been explained by some chemical difference between the D'' layer and the rest of the lower mantle. However, using a combination of dynamical and seismic modelling, Sidorin et al.⁴⁻⁶ have shown that the most consistent explanation is a phase transition in mantle minerals. Their best model^{5,6} had a shear-wave discontinuity of $\sim 1\%$ located $\sim 150 \, \text{km}$ above the core-mantle boundary (depth 2,740 km), with a Clapeyron slope of 6 MPa K^{-1} (though values as large as 10 MPa K^{-1} were almost equally acceptable). The discontinuities of the compressional wave velocities and of the density could not be resolved. The models of Sidorin et al.4-6 were very appealing, but the major problem was that no appropriate phase transition was known at the time. Here we show that MgSiO₃ perovskite undergoes a structural phase transition at the conditions corresponding to the top of the D" layer. The predicted seismic signatures of this transition match the seismological inferences of Sidorin et al.^{4–6}.

A key observation made by Ono *et al.*¹⁴ was that Fe_2O_3 , like MgSiO₃, transforms from the corundum (or ilmenite) to the perovskite structure under pressure. As these authors further found¹⁴, a post-perovskite phase of Fe_2O_3 with a CaIrO₃-type *Cmcm* structure¹⁵ (Fig. 1) is stable above 60 GPa. This has led to the idea that a similar structure could be stable for MgSiO₃ at high pressure.

We explored this idea using *ab initio* simulations based on density functional theory within the local density approximation

Theoretical and experimental evidence for a post-perovskite phase of MgSiO₃ in Earth's D["] layer

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The Earth's lower mantle is believed to be composed mainly of $(Mg,Fe)SiO_3$ perovskite, with lesser amounts of (Mg,Fe)O and $CaSiO_3$ (ref. 1). But it has not been possible to explain many unusual properties of the lowermost ~ 150 km of the mantle (the D" layer) with this mineralogy. Here, using *ab initio* simulations and high-pressure experiments, we show that at pressures and



 $\label{eq:Figure 1} \begin{array}{l} \mbox{Structure of the post-perovskite phase of $MgSiO_3$ (calculated at 120 GPa). SiO_6 octahedra and Mg atoms (spheres) are shown. Similar structures are known for Fe_2O_3, $CalrO_3$, $FeUS_3$, $PbTII_3$, $UScS_3$, $KTml_3$, $AgTaS_3$ and $CalnBr_3$.} \end{array}$

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(LDA) and the generalized gradient approximation¹⁶ (GGA). The calculated enthalpy difference (Fig. 2a) indicates that for pure MgSiO₃ the CaIrO₃-structured phase becomes thermodynamically more stable than perovskite well within the range of lower-mantle pressures: at 83.7 GPa in the LDA or at 98.7 GPa in the GGA. It is well known¹⁷ that usually the LDA underestimates transition press-



Figure 2 Stability of the post-perovskite phase. a, Static enthalpy difference showing a phase transition at 98.7(83.7) GPa in the GGA(LDA). GGA calculations used the allelectron PAW method^{27,28} as implemented in the VASP code²⁹, with $1s^22s^2$ core (radius 2 a.u.) for Mg, $1s^22s^22p^6$ core (radius 1.5 a.u.) for Si, $1s^2$ core (radius 1.52 a.u.) for 0. With a plane wave cut-off of 500 eV and the Brillouin zone sampled by Monkhorst-Pack grids, $4 \times 4 \times 4$ for perovskite and $6 \times 6 \times 4$ for post-perovskite (for the latter we used the primitive cell with 10 atoms), total energy differences and pressure are converged to 0.006 meV per atom and 0.25 GPa, respectively. Structural relaxation was done using conjugate gradients, until the total energy changes were below 10^{-4} eV. The fully converged LDA calculations were done using the ABINIT code³⁰ (ABINIT is a common project of the Université Catholique de Louvain, Corning Inc. and other contributors; http://www.abinit.org). These calculations used Troullier-Martins pseudopotentials, plane waves with a cut-off of 1,088 eV, and the same Monkhorst-Pack grids as in the GGA case. b, Pressure-temperature phase diagram: theory and experiment. Experimental points: perovskite (ref. 13) for mantle composition (KLB-1 peridotite), post-perovskite for pure MgSiO₃. The solid line is based on the static transition pressure (98.7 GPa) and a Clapeyron slope of 9.56 MPa K⁻¹ calculated using density-functional perturbation theory and the GGA, with the ABINIT code³⁰. In these calculations, the dynamical matrices were calculated on $2 \times 2 \times 2$ and $3 \times 3 \times 2$ grids in the Brillouin zone for perovskite and post-perovskite, respectively, using density-functional perturbation theory¹⁸. Interpolating these throughout the Brillouin zone, we calculated phonon frequencies at a very dense reciprocal-space mesh. Errors in the calculated frequencies are within 5 cm⁻¹ (mostly less than 1 cm⁻¹). From the resulting phonon spectra we calculated various thermodynamic properties, including the entropy (S) of each phase and the hightemperature Clapeyron slope $dP/dT = \Delta S/\Delta V$.

ures, whereas the GGA slightly (by a few GPa) overestimates them.

The Clapeyron slope was calculated using density-functional perturbation theory¹⁸ to be 9.85 MPa K⁻¹ within the LDA and 9.56 MPa K⁻¹ within the GGA. These numbers agree with a simple formula for the high-temperature entropy change¹⁹ for transitions without coordination number changes (as in this case): $\Delta S = 3nk_{\rm B}\gamma(\Delta V/V)$, where *n* is the number of atoms in volume *V*, ΔV the volume change at transition, $k_{\rm B}$ the Boltzmann constant, and γ the Grüneisen parameter at the transition (1.2 for perovskite^{20,21}). This formula yields a Clapeyron slope of ~8 MPa K⁻¹.

Following this prediction, we found the new phase experimentally. A sample of pure MgSiO₃ was heated with a laser to overcome potential kinetic effects on possible phase transitions. Experimental details have been described elsewhere^{13,22}. The experimental powder diffraction pattern is shown in Fig. 3; we could index reflections not belonging to platinum, platinum carbide, and rhenium gasket in the *Cmcm* space group with lattice parameters very similar to the theoretical ones (Table 1). This confirms the predicted stability of the post-perovskite phase in pure MgSiO₃. Note that the calculated and experimentally determined stability fields are in excellent agreement with each other (Fig. 2b). At temperatures of the D" layer (~3,000 K) the GGA transition pressure is 127 GPa, which corresponds to the top of the D" layer, 2,740 km depth. We recall that Sidorin *et al.*⁴⁻⁶ suggested a transition at 127 GPa with a Clapeyron slope of 6 MPa K⁻¹.

Equation of state parameters for perovskite and post-perovskite are listed in Table 2, and the elastic constants calculated from stress-



Figure 3 Experimental powder diffraction pattern at 118 GPa and 300 K matching the predicted lattice parameters. The sample was heated with a TEM01-mode YLF laser from two sides without a pressure transmitting medium. The pressure was determined using the reference equation of state of platinum (which was mixed with the sample). The sample temperatures were measured from two sides using the spectroradiometric method. The samples were probed using an angle-dispersive X-ray diffraction technique at the synchrotron beam line BL10XU, SPring-8 in Japan. Filled circles, post-perovskite; Pt, platinum used as an internal pressure calibrant; Re, rhenium of the gasket; PtC, platinum carbide. Observed unit cell dimensions of post-perovskite are a = 2.471(4) Å, b = 8.091(12) Å, c = 6.110(11) Å, V = 122.16(34) Å³. Observed volume change from perovskite to post-perovskite is 0.9%. Vertical bars indicate the calculated positions of the diffraction lines of *Cmcm* post-perovskite phase, *Pbnm* perovskite, platinum, NaCI-type platinum carbide, and rhenium gasket.

strain relations at 120 GPa are given in Table 3. The density discontinuity at the transition is 1.4%; for the mantle the expected density discontinuity is 1.1%. The predicted shear wave discontinuity in MgSiO₃ is 1.9% (1.4% for the mantle), consistent with \sim 1% suggested by Sidorin *et al.*^{4–6}. We predict a very small discontinuity for the compressional wave velocity (0.3% for pure MgSiO₃), explaining why it was so seldom found at the top of the D["] layer^{4–6}.

The calculated properties of our new phase explain many puzzling features of the D" layer. In seismological models, horizontally polarized shear waves are faster (by 1% on average²) than the vertically polarized ones ($v_{SH} > v_{SV}$). Significant seismic anisotropy of the D" layer, containing a signature of the convective flow², could not be explained even qualitatively by mineral physics^{23,24}. The structure of the post-perovskite phase has silicate layers parallel to {010}, which is then the most natural slip plane that will be oriented parallel to the convective flow. In regions of horizontal flow (most of D"), we obtain $v_{SH}/v_{SV} = 1.029 > 1$ for post-perovskite (using the method of ref. 25 and data of Table 3). This is consistent with seismological evidence, and suggests either a large degree of lattice-preferred orientation or significant contributions from other sources such as shape-preferred orientation (anisotropy due to the ordered distribution of crystals and inclusions in the rock). In regions of upwelling convective streams (for example, below the central Pacific), slip planes would be predominantly vertical and one would obtain $v_{SH} < v_{SV}$ actually observed in such regions².

Another mystery of D'', the anticorrelation between the shear (v_s) and bulk sound (v_{ϕ}) velocities^{7,8}, can also be quantitatively explained by the phase transition of MgSiO₃ from perovskite into the post-perovskite phase. We recall that it has been difficult to explain why, at a given depth in the lowermost mantle, anomalies of v_s and v_{ϕ} have opposite signs. As Table 3 shows, the post-perovskite

Table 1 Structures of post-perovskite and perovskite at 120 GPa										
Post-perovskite*				Perovskite†						
Mg Si O1 O2	0 0 0 0	0.2532 0 0.9276 0.6356	1/4 0 1/4 0.4413	Mg Si O1 O2	0.5246 1/4 0.1164 0.1829	0.5768 0 0.4669 0.1926	1/4 1/4 1/4 0.5575			

Table shows GGA results.

Table shows GAA results. * Space group *Cmcm*: a = 2.474 Å, *b* = 8.121 Å, *c* = 6.138 Å; distances (in Å): Mg–O1 = 1.880 (×2), Mg–O2 = 1.955(×4), 2.099(×2); Si–O1 = 1.643 (×2), Si–O2 = 1.695 (×4). † Space group *Phrm*: a = 4.318 Å, b = 4.595 Å, *c* = 6.305 Å; distances (in Å): Mg–O1 = 1.833(×1), 1.893(×1), Mg–O2 = 1.864(×2), 2.047(×2), 2.201(×2); Si–O1 = 1.661(×2), Si–O2 = 1.659(×2), 1.670(×2).

Table 2 Vinet equations of state of perovskite and post-perovskite								
Parameters	Post-pe	erovskite	Pero	Exp.*				
	LDA	GGA	LDA	GGA				
V ₀ (Å ³)	162.86	167.64	163.35	167.42	162.3			
K _ρ (GPa) K _o	231.93 4.430	199.96 4.541	259.82 4.060	230.05 4.142	259.8 3.69			

*Third-order Birch-Murnaghan equation of state parameters at room temperature¹⁰.

transition has a positive jump of v_s and a negative jump of v_{ϕ} . Note that the transition is first order, and in a multicomponent system such as the Earth' mantle there will be an interval ΔT of coexistence of the two phases at given pressure. For this two-phase region, we can write:

$$\left(\frac{\partial \nu_{\phi}}{\partial \nu_{S}}\right)_{p} \approx \frac{\left(\frac{\partial \nu_{\phi}}{\partial T}\right)_{P,x} + f \frac{\nu_{\phi2} - \nu_{\phi1}}{\Delta T}}{\left(\frac{\partial \nu_{\phi}}{\partial T}\right)_{P,x} + f \frac{\nu_{S2} - \nu_{S1}}{\Delta T}}$$
(1)

which includes purely thermal responses (at given composition *x* and pressure *P*) of the velocities and effects due to a phase transition; *f* is the volume fraction of MgSiO₃ (75%). The poorly known effects of variation of Al and Fe content are not included in equation (1), but we find that anticorrelation can be explained without them. Using equation (1), data of Table 3, and assuming thermal responses equal to those of perovskite²⁰, we get $(\partial \ln v_S / \partial \ln v_{\phi})_P = -0.15$ and -0.33 for $\Delta T = 250$ K and 50 K, respectively (from seismic tomography: -0.15 (ref. 8) and -0.3 (ref. 7)). Furthermore, we can reproduce the positive correlation between the shear and compressional (v_P) velocities with the same ΔT : $(\partial \ln v_S / \partial \ln v_P)_P = 3.36$ for $\Delta T = 250$ K (3.3 from ref. 8).

The fact that so many previously unexplained seismic features of the D" layer (seismic discontinuity, its magnitude and Clapeyron slope, anisotropy, bulk-shear velocity anticorrelation) are naturally explained by the properties of post-perovskite is a strong indication that this phase is indeed the major component of the D" layer. The D" layer is not necessarily chemically very different from the rest of the lower mantle, but it surely is different mineralogically. Neither theoretical and experimental error bars (a few GPa for transition pressure) nor the effects of temperature as explored here would change our prediction that the CaIrO₃-type post-perovskite phase is stable in the D" layer. At present, very little is known about the effects of Fe²⁺, Fe³⁺ and Al³⁺ impurities on mantle minerals. We expect that at least Fe³⁺ impurities should stabilize the postperovskite phase against perovskite; this is because at high pressure Fe₂O₃ also has the CaIrO₃ structure¹⁴.

Our findings have other implications. For instance, rheological properties of post-perovskite (probably very different from those of perovskite) and the predicted density discontinuity (1.1%) at top of the D["] layer could be important for mantle dynamics. Also, element partitioning between post-perovskite, perovskite, and molten Fe might be a key to some geochemical anomalies. Elements that are incompatible in the mantle (for example, Na, K, U, Th) might be more easily accommodated in the layered post-perovskite structure, which may affect the chemistry of plume magmas. As Fe₂O₃ at high pressure has the same structure as post-perovskite, post-perovskite could have a greater concentration of Fe³⁺ than perovskite. It is also likely that the size of the D" region increased significantly with time as the mantle cooled down; this is because the Clapeyron slope of the post-perovskite transition is large, $8.0-9.6 \text{ MPa K}^{-1}$. If the whole mantle had been molten with temperatures above 4,000-4,500 K in the early history of the Earth, it would be perovskite that crystallized from the cooling melt, and only on further cooling would post-perovskite and the D" layer have appeared. The presentday thickness of the D" could be used to estimate its age, given the cooling history of the lowermost mantle. As post-perovskite stability requires pressures unattainable in smaller planets like Mercury

Table 3 Elastic constants of perovskite and post-perovskite at 120 GPa											
	C ₁₁	C ₂₂	C ₃₃	C ₁₂	C ₁₃	C ₂₃	C 44	C 55	C 66	K	G
Perovskite*	907	1,157	1,104	513	406	431	364	271	333	648.0	310.9
Post-perovskite†	1,252	929	1,233	414	325	478	277	266	408	647.2	327.5

Table shows GGA results. All elastic constants are in GPa.

*Acoustic velocities (m s⁻¹): $v_P = 14,118$; $v_S = 7,636$; $v_{\phi} = 11,026$. †Acoustic velocities (m s⁻¹): $v_P = 14,158$; $v_S = 7,783$, $v_{\phi} = 10,940$.

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and Mars, many features of these planets would be different from those of the Earth. Further studies are necessary to address these and other issues (elasticity and anelasticity, electrical conductivity, radiative conductivity, energetics of stacking faults, effects of impurities on stability and properties of post-perovskite). Finally, we note that the results of a recent, independent, experimental study²⁶ of the post-perovskite phase transition are consistent with our theoretical and experimental findings.

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Audience drives male songbird response to partner's voice

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According to the social intelligence hypothesis, social context represents an important force driving the selection of animal cognitive abilities such as the capacity to estimate the nature of the social relationships between other individuals¹⁻⁴. Despite this importance, the influence of this force has been assessed only in primates and never in other animals showing social interactions⁵⁻⁷. In this way, avian communication generally takes place in a network of signallers and receivers, which represents an audience altering individual signalling behaviours^{8,9}. Indeed, vocal amplitude¹⁰ and repertoire¹¹ are known to be socially regulated and the attitude towards the opposite sex may change depending on the audience^{8,12,13}. This 'audience effect'^{8,14-16} provides support for the reality of social awareness in some bird species. However no evidence has yet been found to suggest that birds are able to estimate the characteristics of the social relationships between group-mates. Here we show that the male of a gregarious songbird species-the zebra finch (Taeniopygia guttata)-pays attention to the mating status of conspecific pairs, and uses this information to control its behaviour towards its female partner.

Zebra finches are monogamous flock-forming birds that seem to use acoustic recognition for pair-bond maintenance^{11,17}. A number of different vocalizations are produced by this species¹¹, distance calls being the most frequently emitted by both males and females. Distance calls are used by the members of a pair to remain in contact when the flock is foraging or feeding and especially when the birds lose visual contact with each other¹¹. As in many gregarious species, vocal recognition is thus likely to be a key component of reproductive success and it should be supported in both sexes by acoustic cues of distance calls¹⁸⁻²⁰. Previous laboratory experiments testing isolated birds demonstrated that the female zebra finch is able to recognize its mate's vocalizations from other males' vocalizations¹⁷, but never succeeded in demonstrating a mutual acoustic recognition between mates¹¹. In the natural biological context described above, it is very unlikely that wild male zebra finches do not recognize their mates' voices. Two main hypotheses can thus be envisaged: either captive zebra finches have lost some cognitive capacities because of domestication (for instance, females' calls may be less individualized, and/or males may no longer be able to recognize them), or tested males do not show preferential response to their mate's voice owing to a modification of their natural behaviour by their socially isolated position during the playback tests. Indeed social isolation could be a situation of stress in comparison to the natural context where the zebra finch lives in large groups and experiences permanent social interactions that may influence mate-directed behaviour.

To determine whether the vocalizations of female zebra finches support mate recognition, we analysed the acoustic structure of distance calls, searching for acoustic cues which could encode the emitter's individual identity. The female distance call is a complex sound with a fundamental frequency associated with several harmonics (Fig. 1a). This sound is frequency- and amplitudemodulated. With reference to frequency-modulation characteristics, the distance call can be divided into three segments of