

S1 Appendix Details about initial state and construction of unitary matrices used to generate predictions.

The initial state at the beginning of a trial and before a question appears is defined as $\psi(0) = [\psi_i(0)]$, with zeros assigned to all states except for $\frac{1}{\sqrt{11}}$ assigned to the 11 states i = 45, ..., 50, ...55 in the neighborhood of the neutral state corresponding to the middle rating R = 5.

The unitary matrices were constructed from a particular type of quantum random walk model called the Feynman crystal model. Unitary matrices can be constructed from a matrix exponential $U = exp(-i \cdot H)$, where H is a Hamiltonian matrix (a Hermitian matrix). The Hamiltonian matrix we used, $H = [h_{ij}]$, is a tri-diagonal matrix. The entries $h_{i-1,j} = \alpha$ above the diagonal and $h_{i+1,j} = \alpha$ below the diagonal allow diffusion of amplitudes to adjacent states. The entries on the diagonal $h_{ii} = \beta \cdot (\frac{i}{N})$ serve as the potential function. The potential on the diagonal corresponds to a linear potential function that produces constant force in the direction determined by β . The off diagonal entries determine the diffusion rate. We used a Hamiltonian matrix H_A with parameters (α_A, β_A) for the one issue, and we used another Hamiltonian matrix H_B with parameters for each of the two different types of questions.

The best fitting parameters from [?] were used to generate the predictions. We set $\beta = -14.5725$, $\alpha = 4.5975$ for issue A, and we set $\alpha = 4.5975$ for issue B and varied β from -30 to +30 to sweep out the model predictions shown in Figure 1. The middle of the range for β on issue B corresponds to the value of β used for issue A, and so the questions become compatible at this point.

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