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Ro-vibrational excitation of H_2 by H for high temperature regime



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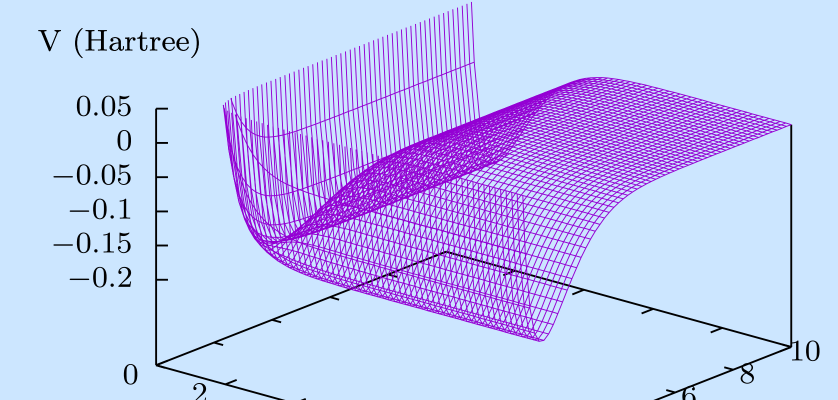
Context : $H+H_2$ collisions

Accurate knowledge of the abundance of H_2 molecules is crucial for the thermal balance of astrophysical media (early Universe, post-shock regions,...). Need to study the main collisions leading to the formation/destruction of H_2 molecules for temperature ranging from few Kelvins (interstellar medium) to high temperatures (15 000 K). There is low to moderate (5000 K) temperature regime studies using quantum mechanical approaches (TIQM)[1] with an accurate H₃ potential energy surface [3].

Objectives : Study the effects of high energy collisions of H_2 by H compared to low to moderate temperature regimes. Molecular hydrogen is ortho hydrogen (nuclear spin $\uparrow\uparrow$: odd rotational quantum numbers) or para hydrogen (nuclear spin $\uparrow\downarrow$: even rotational quantum numbers).

 $H_A + H_B H_C(v', j')$ inelastic, $H_A + H_B H_C(v, j) \rightarrow \begin{cases} H_B + H_A H_C(v', j') & reactive, \end{cases}$ $\begin{array}{c} \mathbf{H}_{C} + \mathbf{H}_{A}\mathbf{H}_{B}(v',j') \\ \mathbf{H}_{A} + \mathbf{H}_{B} + \mathbf{H}_{C} \end{array}$ reactive,

Ortho-para-H₂ ratio important for the cooling mechanism of H_2 , **reactive collision** is the only process for the ortho-para- H_2 conversion.



 $\mathbf{r}_{BC}(a_0)$ $\mathbf{R}_{A,BC}(a_0)^6$

Quasi-Classical Trajectory (QCT) calculation

dissociative,

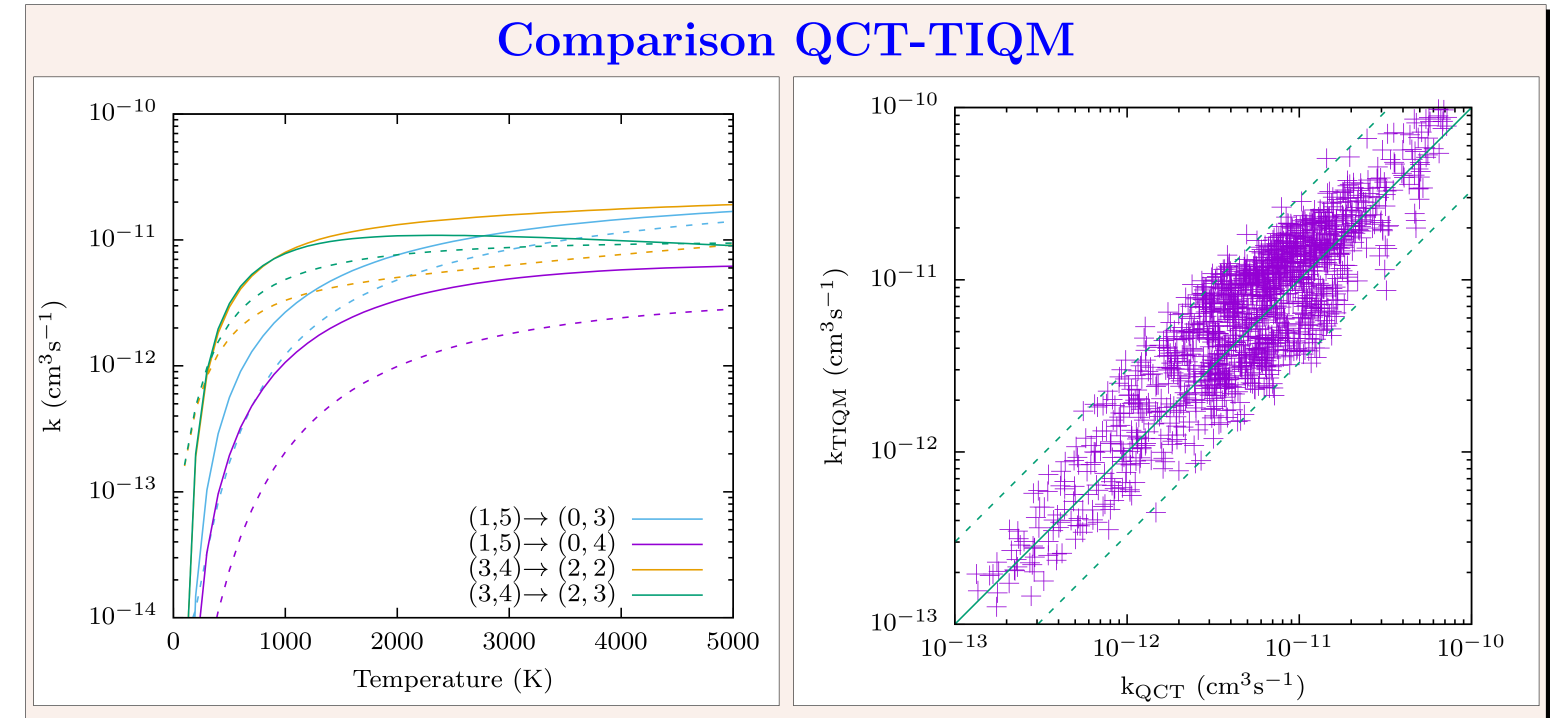
- State-to-state calculations on our own QCT code,
- Initial ro-vibrational quantum states (v, j) with energies ranging from 0 to 50 000 K,
- Nuclear dynamics solving Hamilton's equations of motion,
- Collisional energies E_{col} ranging from 0.025 Hartree (0.068 eV) to 0.18 Hartree (4.9 eV) every 0.005 Hartree (0.136 eV),
- Assignment of final ro-vibrational state using a standard binning method[4],
- All final ro-vibrational bounded states (v', j') taken into account,
- Cross section function of the reactive probability P_r :

 $P_r(v, j \to v', j'; b, E_{col}) = \frac{N_r(v', j'; b, E_{col})}{N(v, j; b, E_{col})};$ $\sigma_{v', j' \leftarrow v, j}(E_{col}) = \pi b_{max}^2 P_r(v, j \to v', j'; E_{col})$

• State-to-state rate constant :

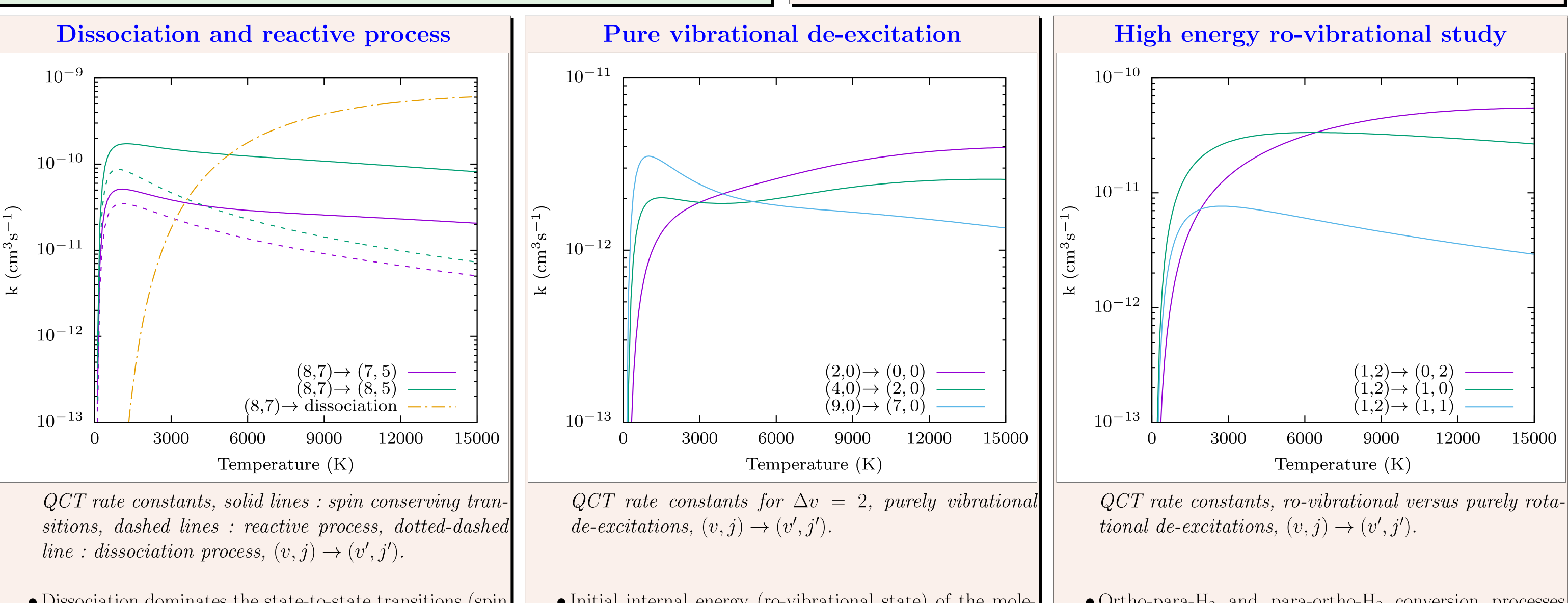
$$k_{v',j'\leftarrow v,j}(T) = \left(\frac{8k_BT}{\pi\mu_{A,BC}}\right)^{1/2} \frac{1}{(k_BT)^2} \int_0^\infty \sigma_{v',j'\leftarrow v,j}(E_{col}) E_{col} \exp\left(-\frac{E_{col}}{k_BT}\right) dE_{col}$$

b impact parameter, N_r number of reactive trajectories, N total number of trajectories, k_B boltzman constant and $\mu_{A,BC}$ the reduced mass of the system.



Left : rate constants results for QCT (solid lines) and TIQM (dashed lines), $(v, j) \rightarrow$ (v', j'); right : comparison between QCT and TIQM rate constants at 5000 K.

• Agreement between TIQM[1] and QCT calculations below a factor of 3 for spin conversion process (reactive collisions) as for spin conservation process (inelastic + reactive collisions). • Allows to extend the QCT calculations above the energies reachable using quantum methods.



• Dissociation dominates the state-to-state transitions (spin

• Initial internal energy (ro-vibrational state) of the mole-

• Ortho-para- H_2 and para-ortho- H_2 conversion processes

- conservation and conversion processes) above few thousands of Kelvins for the collisions involving quantum states of internal energies above 3.5 eV,
- De-excitation rate constants increase with the internal energy of the final quantum state,
- Rate constant for the reactive process dominates the collisions around 1000 to 3000 K,
- Inelastic process dominates for low total energy collisions (low collisional energy and/or low initial state energy) and also for the high temperature regime (high collisional energy and/or high initial state energy).

cule contributes the most to the rate constant at low to moderate temperature collisions,

- Collisional energy leads the rate constant at high temperature (compared to internal energy),
- Leads to a crossing between high internal energy rate constants and high collisional energy rate constants,
- Concerning high internal energy states $(\text{see } (4,0) \rightarrow (2,0))$: inelastic collisions dominates high temperature (with dissociation) \rightarrow minimum of the rate constant around 5000 K,

• Similar for ro-vibrational de-excitations (for small Δj).

(only reactive collisions) always slower than spin conservation processes (reactive + inelastic collisions) as expected[2] for all the temperature range considered, • Purely rotational de-excitations faster than ro-vibrational de-excitations for low collisional energy collisions, • For high collisional energy collisions (lot of energy to store into internal energy) as $\Delta E_v > \Delta E_i$, ro-vibrational deexcitations faster,

• Leads to a crossing between purely rotational de-excitation and ro-vibrational de-excitation rate constants moving to higher temperature with higher initial internal energy.

References :

1. F. Lique, Mon. Not. R. Astron. Soc., 453, 1 (2015) 2. S.L. Mielke, B.C. Garrett, K.A. Peterson, J. Chem. Phys., 116, 4142 (2002) 3. D.G. Truhlar, J.T. Muckermann in R.B. Bernstein ed., Atom-Molecule Collision Theory, 4. D. Bossion, Y. Scribano, F. Lique, G. Parlant, Mon. Not. R. Astron. Soc., submitted.

Conclusion

- Low to moderate temperature rate constants behavior should not be extrapolated for high temperature regime due to a non-monotonic evolution of the state-to-state rate constants,
- Dissociation process cannot be neglected for the high temperature regime [4].