

Study of atomic motion in rubidium borate glasses

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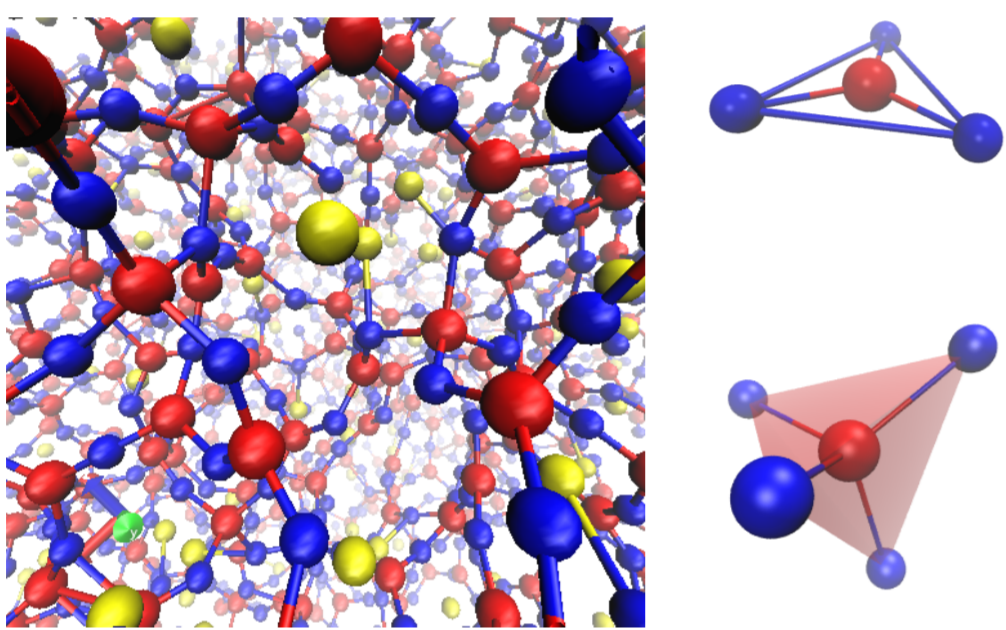
Introduction

Atomic Scale X-ray Photon Correlation Spectroscopy (aXPCS) uses coherent X-rays to probe the dynamics of materials on an atomic scale. It was applied to study atomic diffusion in crystals [1], but its application was

recently extended to glasses as well [2]. Results of dynamics and Small-Angle X-ray Scattering (SAXS) studies on high- and low-alkali content rubidium borate glasses are presented here.

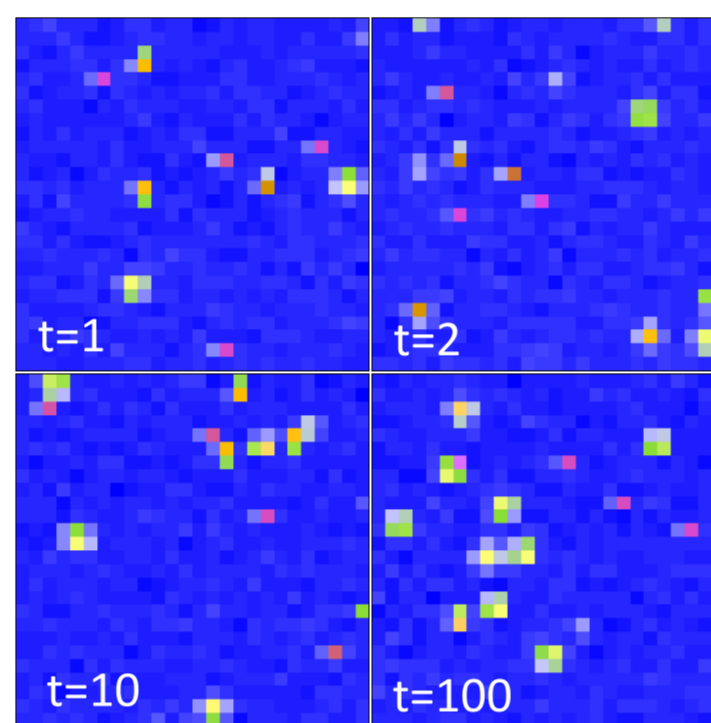
Theory

Real space structure: changes in time results in different scattering patterns



Space and time dependence described by the Van Hove Pair Correlation Function

Sequence of scattering patterns
→ Time correlation

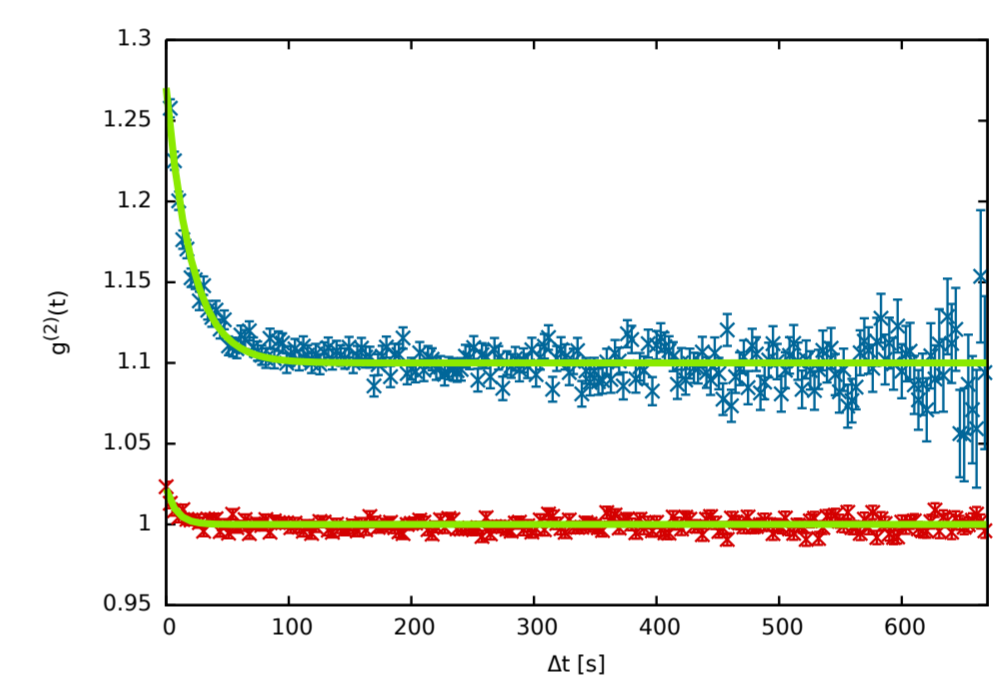


Intensity Auto Correlation Function (ACF)

$$g^{(2)}(\vec{q}, \Delta t) := \frac{\langle I(\vec{q}, t)I(\vec{q}, t + \Delta t) \rangle}{\langle I(\vec{q}, t) \rangle^2}$$

Functional form of the ACF:

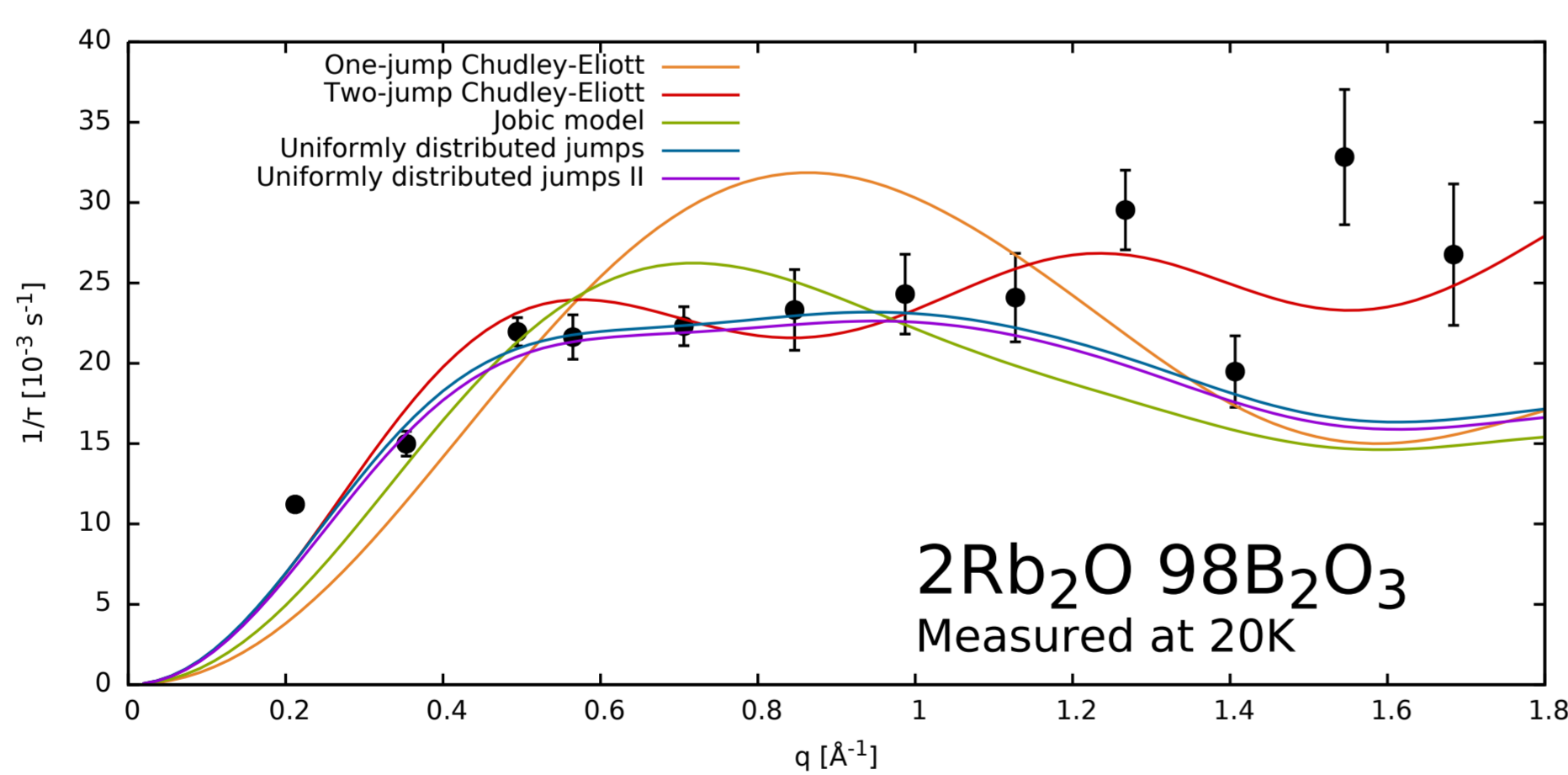
$$g^{(2)}(q) = 1 + \beta e^{-\left(\frac{2\Delta t}{\tau}\right)^\alpha}$$



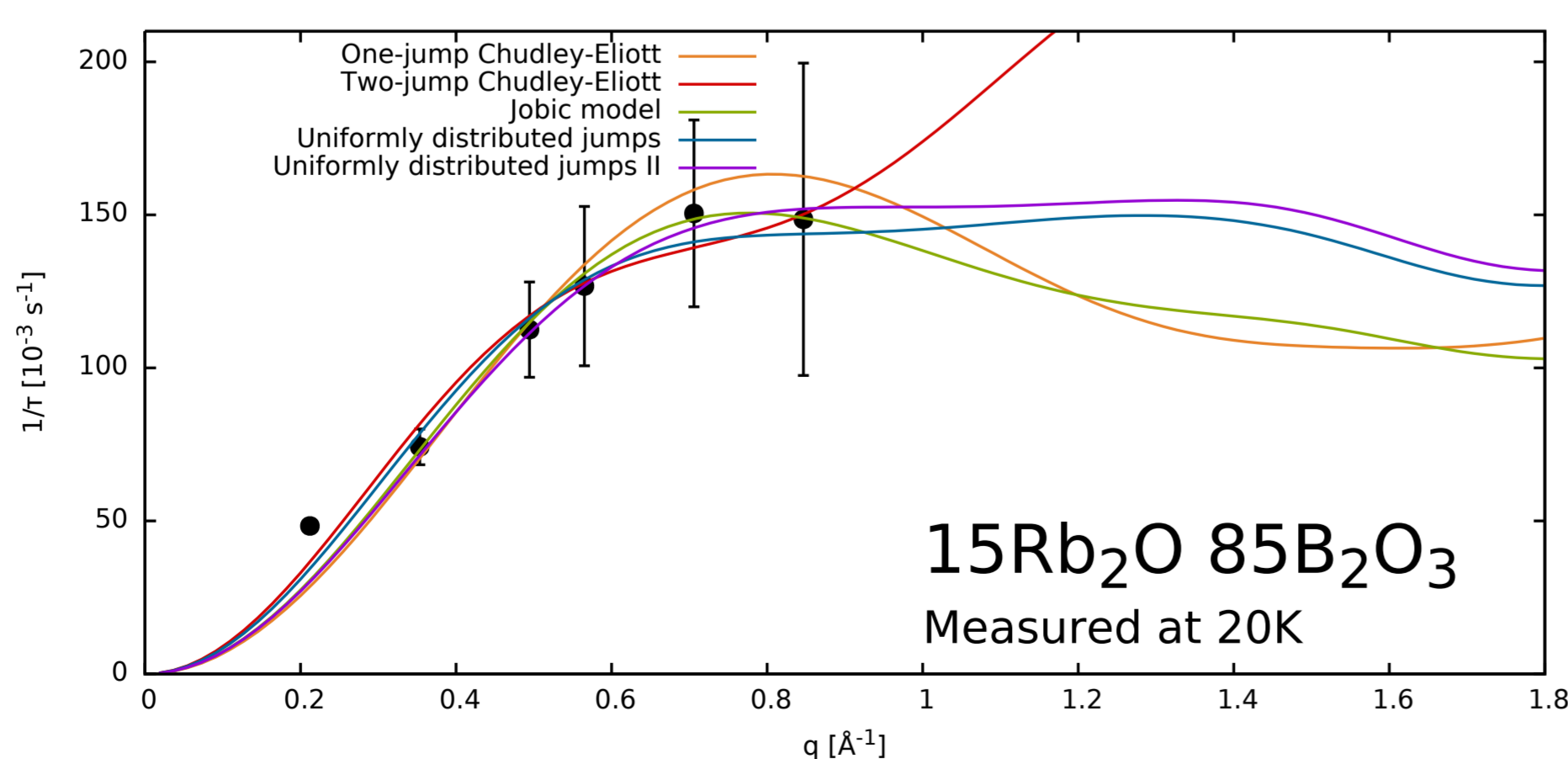
Normalization by the total structure factor

$$\tau_{coh}(\vec{q})^{-1} = \frac{\tau_{inc}(\vec{q})^{-1}}{S_{total}(q)}$$

Results: fit to jump models

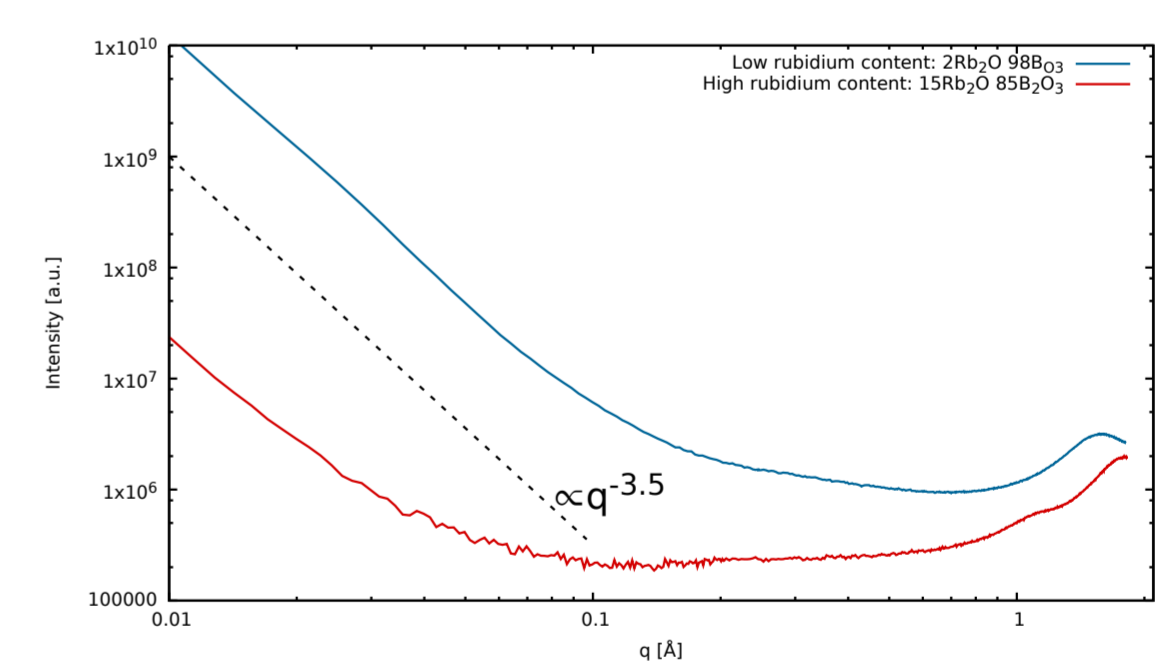


Best fit:
Two-jump
distances
Chudley-Eliott
Model with
jump lengths
 $d_1 = 1.0 \text{ \AA}$ and
 $d_2 = 8.5 \text{ \AA}$, with
3/4 of
short-distance
jumps



top: low alkali
content glass,
bottom: High
alkali content
glass. Several
models can be
fitted at small
q-range. Fits
are equally
good in the
hydrodynamic
limit.

SAXS



SAXS intensities as a function of momentum transfer

Conclusion

- Diffusion in low rubidium content borate glasses best described by two-jump model
- Large difference in diffusivity between low and high alkali glass
- Diffusion at 20K still too fast to be measured in high alkali glass
- Inhomogeneities are much larger in high alkali sample