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Two inherent crossovers of diffusion process in glass-forming liquids

Maiko Kofu,^{1,*} Antonio Faraone,^{2,3} Madhusudan Tyagi,^{2,3} Michihiro Nagao,^{2,4} and Osamu Yamamuro^{1,†}

¹Institute for Solid State Physics, University of Tokyo, Kashiwa, Chiba 277-8581, Japan

²NIST Center for Neutron Research, National Institute of Standards and Technology, Gaithersburg, Maryland 20899-6102, USA

³Department of Materials Science, University of Maryland, College Park, Maryland 20742, USA

⁴Center for Exploration of Energy and Matter, Indiana University, Bloomington, Indiana 47408-1398, USA

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We report on incoherent quasielastic neutron scattering measurements examining a self-diffusion process in two types of glass-forming liquids, a molecular liquid (3-methylpentane) and an ionic liquid (1-butyl-3-methylimidazolium bis(trifluoromethanesulfonyl)imide). We have experimentally demonstrated that both liquids exhibit two crossovers in the momentum transfer (Q) dependence of the self-correlation function which is basically described by the stretched exponential function, $\exp[-(t/\tau)^{\beta}]$. The first crossover point ($Q \approx 0.2 \text{ Å}^{-1}$) corresponds to a crossover from Fickian ($\beta = 1$) to non-Fckian ($\beta \neq 1$) diffusion attributed to dynamical correlation. On the other hand, the second one at $Q \approx 0.8 \text{ Å}^{-1}$ is associated with the crossover from the Gaussian to non-Gaussian behavior. It is remarkable that the stretching exponent β gradually changes in between the two crossover points. We consider that the two crossovers are the universal feature for glass-forming liquids.

I. INTRODUCTION

It has been believed that the self-diffusion of molecules 9 ¹⁰ in a liquid is basically governed by Fick's law. The self-¹¹ part of the van Hove correlation function $G_{\rm s}(r,t)$ is writ-¹² ten as the Gaussian function with respect to r and de- $_{13}$ cays exponentially with t (Debye relaxation). Incoher-¹⁴ ent quasielastic neutron scattering (IQENS) experimen-¹⁵ tally gives the Fourier transform of $G_{\rm s}(r,t)$, the inco-¹⁶ herent dynamical structure factor $S_i(Q,\omega)$ or the inco-¹⁷ herent intermediate scattering function $I_{\rm s}(Q,t)$, where ¹⁸ Q and ω are the momentum and energy transfer. In ¹⁹ the case of the Fickian diffusion, the relaxation obeys ²⁰ the relation $I_{\rm s}(Q,t) = \exp(-t/\tau) = \exp(-DQ^2t)$, where $_{21}$ D denotes the self-diffusion coefficient and τ is the re-22 laxation time. On the other hand, the diffusion pro-23 cess in supercooled liquids is typically characterized ²⁴ by the non-Debye KWW (Kohlrausch-Williams-Watts) ²⁵ function, $\exp[-(t/\tau)^{\beta}]$, where the exponent β is a stretching parameter accounting for the deviations from the sim-26 $_{27}$ ple exponential behavior. A natural question arises as to $_{28}$ whether the relation $\tau\propto Q^{-2}$ is valid for the non-Debye ²⁹ diffusion process. Many attempts to understand the na-³⁰ ture of dynamics in the supercooled regime, including $_{31}$ its Q-dependence, have been made by theoretical calcu-₃₂ lations/simulations [1–19] and guasielastic neutron scattering experiments [20–33]. 33

³⁴ A crossover in the *Q*-dependence of self-correlation ³⁵ functions was first discussed in glass forming poly-³⁶ mers [11, 14, 20–30]. Both IQENS and molecular dy-³⁷ namics (MD) simulations have shown that the relaxation ³⁸ time follows the relation $\tau(Q) \propto Q^{-2/\beta}$ below Q_2 while ³⁹ $\tau(Q) \propto Q^{-2}$ above Q_2 , where Q_2 roughly corresponds



FIG. 1. Molecular (Ionic) structures of (a) 3methylpentane (3MP) and (b) 1-butyl-3-methylimidazolium bis(trifluoromethanesulfonyl)imide (C4mimTFSI).

⁴⁰ to the first maximum of the static structure factor S(Q). ⁴¹ The change at Q_2 is interpreted as a crossover from Gaus-⁴² sian at lower Q to non-Gaussian nature at higher Q. Ad-⁴³ ditional crossover, which occurs at Q_1 ($< Q_2$), is reported ⁴⁴ for supercooled glycerol [31, 33] and monomer species of ⁴⁵ polymer [17]. The relaxation time is proportional to as ⁴⁶ Q^{-2} below Q_1 , which is expected for the ordinary Fick-⁴⁷ ian diffusion. Note that the absence of the crossover at ⁴⁸ Q_1 in polymers is a consequence of the chain connectiv-⁴⁹ ity of macromolecules giving rise to a different type of ⁵⁰ dynamics, Rouse dynamics.

The two crossovers have so far been found by MD s2 simulations and fragmentally observed in IQENS exs3 periments [17, 31, 33]. Only in glycerol, the detailed Q-dependence of τ and β are investigated experimens5 tally [31]. However, $I_{\rm s}(Q, t)$ are plotted against reduced times estimated assuming the scaling of the characteriss7 tic time τ , viscosity η , and temperature ($\tau \propto \eta(T)/T$); se the data taken at different temperatures are combined to s9 make a single master $I_{\rm s}(Q, t)$ curve covering the whole Qso region. Therefore there remains serious ambiguity in the s1 spatial scale dependence of diffusion behavior, especially s2 in the variation of β .

In this paper, we demonstrate that $I_{\rm s}(Q,t)$ ex-

^{*} Present address: Materials and Life Science Division, J-PARC Center, Tokai, Ibaraki 319-1195, Japan; maiko.kofu@j-parc.jp

[†] yamamuro@issp.u-tokyo.ac.jp

64 hibits the two crossovers in two different types 65 of liquids, a molecular liquid (3-methylpentane) 66 and an ionic liquid (1-butyl-3-methylimidazolium 67 bis(trifluoromethanesulfonyl)imide, C4mimTFSI), using 68 real experimental data. Molecular structures of both 69 liquids are shown in Fig. 1. The glass transition tem-⁷⁰ peratures (T_g) are reported to be 77 K for 3MP [34] and ⁷¹ 181.5 K for C4mimTFSI [35]. There are several types 72 of glasses including network glass, molten salt glass, 73 molecular glass, hydrogen-bond glass, metallic glass, and ⁷⁴ polymer glass. Glycerol is an example of hydrogen bond ⁷⁵ glass and 3MP is a typical molecular glass. Ionic liquids 76 are intermediate between molecular and molten salt 77 glasses, owing to their amphipathic nature. The study 78 on 3MP and C4mimTFSI, including the previous results ⁷⁹ on glycerol, allows us to clarify whether the crossover is ⁸⁰ a generic feature in the diffusion process of glass-forming 81 liquids.

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II. EXPERIMENTAL

3MP and C4mimTFSI, whose purities were better than 83 84 99.9 %, were purchased from Tokyo Chemical Industry 85 Co., Ltd. and Kanto Chemical Co., Inc. respectively ⁸⁶ and used without further purification. IQENS experi-⁸⁷ ments were made on a neutron spin echo spectrometer ⁸⁸ NSE [36, 37], a backscattering spectrometer HFBS [38] ⁸⁹ and a disk chopper time-of-flight spectrometer DCS [39] ⁹⁰ at the NIST Center for Neutron Research (NCNR) of the National Institute of Standards and Technology (NIST) 124 91 ⁹² in the USA. For 3MP and C4mimTFSI, the observed ¹²⁵ HFBS instrument, to obtain the mean-square displacescattering signal is dominated by the incoherent scatter- $_{126}$ ment $\langle u^2 \rangle$ in a wide temperature range. $\langle u^2 \rangle$ is evalu-⁹⁴ ing from H atoms and so we measure the time evolution ¹²⁷ ated from the equation, $S(Q,\omega) \propto \exp(-\langle u^2 \rangle Q^2/3)$. The ⁹⁵ of self diffusion of tagged particles (H atoms).

 $_{97}$ tron wavelengths (λ) of 4.5 Å, 6 Å, and 8 Å with a distri- $_{130}$ peratures the dynamics enters the time window of the ⁹⁸ bution of $\Delta\lambda/\lambda = 0.15$ at a total Q range from 0.15 Å⁻¹ ¹³¹ neutron spectrometer. The $\langle u^2 \rangle$ data between 213 K and ³⁰ ballon \hat{A}^{-1} . The range of Fourier times was between 4 ps ¹³² 240 K for C4mimTFSI are missing due to crystallization. ¹⁰⁰ and 35 ns. The HFBS instrument was operated in both ¹³³ At around $T_{\rm g}$, $\langle u^2 \rangle$ displays a clear upturn, indicating ¹⁰¹ the fixed window and dynamic window modes. In the ¹³⁴ that some relaxation gets activated. It should be empha-¹⁰² dynamic window mode for IQENS experiments, Doppler ¹³⁵ sized that $\langle u^2 \rangle$ of 3MP steeply increases above $T \simeq 1.3 T_{\rm g}$ ¹⁰³ shifted neutrons with $\lambda = 6.27$ Å allow the investigation ¹³⁶ while that of C4mimTFSI gradually increases. The re- $_{104}$ of the dynamic range of \pm 15 μ eV with an energy reso- $_{137}$ sults are indicative of multiple relaxation processes in ¹⁰⁵ lution of 0.8 μ eV (full width at half maximum). The Q_{-138} C4mimTFSI and a single process in 3MP. ¹⁰⁶ range covered by HFBS was 0.25 Å⁻¹ $\leq Q \leq 1.75$ Å⁻¹. ¹³⁹ Figure 3 shows the normalized intermediate scatter- $_{107}$ As for C4mimTFSI having fast local motions, the IQENS $_{140}$ ing functions, $I_{\rm s}(Q,t)/I_{\rm s}(Q,0)$, at several Qs between ¹⁰⁸ measurement was also made on DCS utilizing incident ¹⁴¹ 0.15 Å⁻¹ and 1.0 Å⁻¹. The data measured on two or ¹⁰⁹ neutrons of 6 Å and 9 Å with corresponding resolutions ¹⁴² three spectrometers are combined to produce the real ¹¹⁰ of 64 μ eV and 22 μ eV and maximum Qs of 1.3 Å⁻¹ and ¹⁴³ $I_s(Q, t)$ curves in the time range of four orders of mag-111 ¹¹² to investigate the diffusion dynamics in wide time (1 ps ¹⁴⁵ values among different spectrometers with different en-¹¹³ to 100 ns) and Q (0.15 Å⁻¹ to 1.7 Å⁻¹) ranges. The ¹⁴⁶ ergy windows. This procedure is often required, because ¹¹⁴ data were corrected at 127 K for 3MP and at 300 K for $_{147}$ $I_{\rm s}(Q,0)$ is practically obtained by the integration over the $_{115}$ C4mimTFSI, which correspond to 1.65 $T_{\rm g}$. The tempera- $_{148}$ finite energy region that is specific to the spectrometers. ¹¹⁶ tures were chosen to examine the Q-dependence of diffu-¹⁴⁹ No scale factor was applied for the data of NSE and DCS $_{117}$ sion dynamics in the accessible time range by the neutron $_{150}$ with $\lambda = 6$ A, where the integral range is about ± 1 meV. ¹¹⁸ spectrometers used. The data were also recorded below ¹⁵¹ While, the scale factors larger than 1 were multiplied for



FIG. 2. (color online) Mean-square displacements $\langle u^2 \rangle$ of 3MP and C4mimTFSI as a function of $T/T_{\rm g}$ taken on HFBS. Temperatures of glass transitions (T_g) and IQENS measurements (T_{meas}) are denoted. The inset shows temperature dependence of $\langle u^2 \rangle$ and the dashed line displays an expectation for a harmonic oscillator.

¹¹⁹ 10 K to obtain the instrumental resolutions. The data ¹²⁰ reduction and Fourier transform of $S_i(Q,\omega)$ data were ¹²¹ performed using the DAVE software package [40] and we ¹²² discuss only $I_{\rm s}(Q,t)$ in this paper.

III. **RESULTS AND DISCUSSION**

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Elastic fixed window scans were performed first on the ¹²⁸ temperature dependence of $\langle u^2 \rangle$ for both liquids are pre-The measurements on NSE were performed using neu- 129 sented in Fig. 2. This plot is helpful to see at what tem-

1.9 Å⁻¹. The use of these three spectrometers enabled us 144 nitude. Note that scale factors are applied to match the



FIG. 3. (color online) Normalized intermediate scattering functions of (a) 3MP at 127 K and (b) C4mimTFSI at 300 K for several Q values from 0.15 Å⁻¹ to 1.0 Å⁻¹.

¹⁵² the other data with narrower integral ranges. In both ¹⁵³ liquids, clear relaxation processes were observed. The $_{154}$ self-correlation function decays faster at higher Q, which ¹⁵⁵ is characteristic of the diffusion process. Apparently, the ¹⁵⁶ shape of the self-correlation function varies depending on $_{157}$ Q; the higher Q is, the more gradually it decays in a wider 158 time range.

The self-correlation functions of 3MP were well fitted 159 ¹⁶⁰ to a single KWW function,

$$\frac{I(Q,t)}{I(Q,0)} = Ae^{-\left(\frac{t}{\tau_{\rm KWW}}\right)^{\beta}}.$$
(1)

¹⁶² damping of vibration and any other local relaxation ²¹⁷ 0.8 Å⁻¹ for 3MP and C4mimTFSI at $T = 1.65 T_g$. This ¹⁶³ hardly occur at a temperature as low as 127 K. One may ²¹⁸ could be due to the relative proximity of Q_1 to Q_2 . We ¹⁶⁴ consider that rotations of CH₃ group could be observed ²¹⁹ anticipate that, as the system is further cooled, the first $_{165}$ even at low temperatures. However, in the present data $_{220}$ crossover occurs at a lower Q and the $\tau \propto Q^{-2/\beta}$ relation $_{166}$ of 3MP, the CH₃ rotations were not independently ob- $_{221}$ is confirmed in a wider Q region. $_{167}$ served from the translation diffusion. The absence of $_{222}$ Figure 4 also presents the static structure factor S(Q)166 CH₃ rotations is also supported by the temperature de- 223 of fully-deuterated 3MP obtained at 140 K with a neu-¹⁶⁹ pendence of $\langle u^2 \rangle$ (Fig. 2); there is no sign of relaxation ²²⁴ tron scattering spectrometer SWAN at KENS, High En- $_{170}$ process below 1.3 $T_{\rm g}$. Perhaps, inter- and intra-molecular $_{225}$ ergy Accelerator Research Organization, Japan, and the $_{171}$ interactions in 3MP can cause high potential barriers for $_{226}$ S(Q) of C4mimTFSI at room temperature with an X-ray ¹⁷² the rotations. On the other hand, local relaxation pro- $_{227}$ diffraction instrument [43]. The first maximum in S(Q), ¹⁷³ cesses were clearly observed in C4mimTFSI. The relax- ²²⁸ Q_{max} , is situated at 1.4 Å⁻¹ for 3MP and 0.85 Å⁻¹ [44]

¹⁷⁵ nation of exponential and KWW functions,

$$\frac{I(Q,t)}{I(Q,0)} = A_0 e^{-\frac{t}{\tau_0}} + A_1 e^{-\frac{t}{\tau_1}} + A_2 e^{-\left(\frac{t}{\tau_{\rm KWW}}\right)^{\beta}}.$$
 (2)

¹⁷⁶ The two exponential terms are attributed to fast local processes and their relaxation times were fixed to the ¹⁷⁸ literature values ($\tau_0 = 0.66 \text{ ps}, \tau_1 = 8.97 \text{ ps}$) [41]. A_i $_{179}$ is the fraction of each relaxation and a constraint, A_0 + $_{180} A_1 + A_2 = \exp(-2W)$, was applied. Here, $\exp(-2W)$ ¹⁸¹ is known as the Debye-Waller factor and was estimated ¹⁸² from the elastic scattering signals taken on DCS.

The product of the mean relaxation time and Q^2 , $_{184} \langle \tau \rangle Q^2$, and β are plotted against Q in Fig. 4. The 185 mean relaxation time is evaluated using the relation 186 $\langle \tau \rangle = \tau_{\rm KWW} / \beta \cdot \Gamma(1/\beta)$ (Γ is the Gamma function). Note 187 that $\langle \tau \rangle Q^2$ is constant (=1/D) when the Fickian diffu-188 sion dynamics is observed. It is evident that the character ¹⁸⁹ of tagged particle motion changes at $Q = Q_1 ~(\approx 0.2 ~\text{\AA}^{-1})$ ¹⁹⁰ and $Q_2 \ (\approx 0.8 \ \text{\AA}^{-1})$ in both liquids. Interestingly, β also ¹⁹¹ changes at these Qs. Below $Q_1, \langle \tau \rangle Q^2$ is almost constant (red lines in Fig. 4) and β is approximately 1. Although the data of C4mimTFSI fluctuate due to the presence of the fast local relaxation processes, the data are in rough ¹⁹⁵ agreement with the value of D^{-1} obtained from a pulsed ¹⁹⁶ field gradient nuclear magnetic resonance (PFG-NMR) ¹⁹⁷ experiment $(Q < 2 \times 10^{-4} \text{ Å}^{-1})$ [42] (dash-dotted red ¹⁹⁸ line). Therefore, it appears reasonable to conclude that ¹⁹⁹ the process below Q_1 is the Fickian diffusion. In the in-²⁰⁰ termediate region ($Q_1 < Q < Q_2$), the Q-dependence of ²⁰¹ $\langle \tau \rangle$ is steeper than Q^{-2} . $I_s(Q,t)$ is not reproduced by ²⁰² the exponential function but the KWW one. The ex-²⁰³ ponent β decreases with increasing Q and reaches 0.5 ²⁰⁴ at Q_2 . Above Q_2 , the relaxation times again follow the ²⁰⁵ relation $\tau \propto Q^{-2}$ (green dashed line) but β remains al- $_{206}$ most constant at 0.5. The variations of τ and β with Q207 are qualitatively similar among 3MP, C4mimTFSI and ²⁰⁸ glycerol [31], though the glycerol data have ambiguity 209 caused by the reduced time. These results suggest that ²¹⁰ the appearance of the two crossovers is generic behavior in supercooled liquids.

Dashed blue curves in Fig. 4 show the relation $\tau \propto$ $_{213} Q^{-2/\beta_{av}}$ implying the Gaussian nature, which is reported ²¹⁴ in previous works [11, 14, 17, 20–31, 33]. Here the average ²¹⁵ values of the stretching parameter, $\beta_{av} = 0.65$, were used. ¹⁶¹ The obtained prefactor A was roughly 1, indicating that ²¹⁶ The relation holds in a narrow Q region 0.4 Å⁻¹ < Q <

 $_{174}$ ation curves of C4mimTFSI were fitted with the combi- $_{229}$ for C4mimTFSI. The crossover occurs at a similar Q_2



FIG. 4. (color online) Mean relaxation times multiplied by Q^2 and the exponents β of KWW functions against Q for (a) 3MP at 127 K and (b) C4mimTFSI at 300 K. Static structure factors S(Q) are also shown as gray shaded areas. In the upper panels, horizontal lines show $\beta = 0.5$ and solid curves are guides for the eyes. In the lower panels, dashed red and occur and Q_{max} at the first maximum in S(Q).

 $_{231}$ ferent. The length scale of Q_2 is about 8 Å and roughly $_{269}$ increased), the memory effect fades away, which causes $_{232}$ corresponds to the average intermolecular distance. In $_{270}$ the increase in β toward 1. Below Q_1 with a length scale $_{233}$ regard to the position of Q_2 , our result is consistent with $_{271}$ larger than 30 Å (blue process), the particle dynamics is $_{234}$ those reported previously; the crossover point Q_2 is lo- $_{272}$ highly coarse-grained in time and resembles the ordinary $_{235}$ cated in the range of 0.6 Å⁻¹ to 1.0 Å⁻¹ and close to but $_{273}$ Fickian diffusion. It should be noted here that the mem- $_{236}$ slightly lower than Q_{max} in polymers [11, 14, 20–30] and $_{274}$ ory effect is convoluted in the diffusion dynamics, even if ²³⁷ glycerol [31, 33].

239 tected in the time and spatial scales of neutron scattering. 277 than that of the simple Fickian diffusion. 240 In glass-forming liquids, even at temperatures well above 278



FIG. 5. (color online) Schematic illustration of diffusion process in short (red; $Q > Q_2$), medium (green; $Q_1 < Q < Q_2$), and long (blue; $Q < Q_1$) spatial scales.

 $_{\rm ^{241}}$ $T_{\rm g}$ (T $\,=\,1.65$ $T_{\rm g}$ in the present case), molecules corre-242 late with each other and they move cooperatively. The ²⁴³ dynamically correlated region is rather small and its size is about the intermolecular distance in the temperature ²⁴⁵ range. The structural correlation causes the sluggish cooperative motion, where memory effects play an essential 247 role in the dynamics. Such a motion is characterized by the KWW function. 248

In IQENS experiments, τ at a given Q corresponds to 249 an average time for a tagged particle stepping out of the 250 region with a size of $2\pi/Q$ (see Fig. 5). Therefore, both 251 crossovers could be discussed in terms of "spatial averag- $_{253}$ ing". In the Q region above Q_2 , the corresponding spatial ²⁵⁴ region is rather small and the tagged particle experiences ²⁵⁵ only one or two jumps to leave the region (red process in Fig. 5). In the case, the particles still remember where 256 ²⁵⁷ they were in the recent past due to the short-range corre-²⁵⁸ lations. The motions of tagged particles inside the corre-²⁵⁹ lated region would be rather heterogeneous, being associ-²⁶⁰ ated with the non-Gaussian behavior. When we take avgreen lines represent the $\tau \propto Q^2$ behavior and dashed blue $_{261}$ erage of the motions over the space larger than the corre-curves denote $\tau \propto Q^{2/\beta_{av}}$ ($\beta_{av} = 0.65$). Dash-dotted red line $_{262}$ lated region ($Q_1 < Q < Q_2$; green process), all of the par-corresponds to D^{-1} estimated from the PFG-NMR experi- $_{263}$ ticles relax identically (Gaussian nature) but intrinsically ment [42]. Q_1 and Q_2 are the Q positions at which crossover $_{264}$ exhibit the KWW behavior due to the memory effect. In ²⁶⁵ this transient region, the particle undergoes a sequence ²⁶⁶ of jumps including back-scattering events, which is often ²⁶⁷ called sub-diffusion, giving rise to the $\tau \propto Q^{-n}$ (n > 2) $_{230}$ ($\approx 0.8 \text{ Å}^{-1}$) in both liquids but Q_{max} is somewhat dif- $_{268}$ behavior. As Q is further decreased (the spatial region is 275 the dynamics seems "uncorrelated". The effective diffu-We now discuss microscopic molecular motions de- 276 sion coefficient with the memory effect could be smaller

The previous MD simulation works have shown that

 $_{260}$ cant value above Q_2 [11, 17, 25–27, 30, 33]. Although $_{328}$ eters with different time windows were used to obtain ²⁸¹ NGP is not estimated from our experiments, it is reason-³²⁹ self-correlation functions in a wide time range. It was 282 able to consider that the Gaussianity changes at around 330 demonstrated that the diffusion behavior of molecules 283 Q₂. Colmenero, Arbe, and co-workers suggested that the 331 changes at two crossover points. Below the first crossover ²⁸⁴ crossover could also be understood as a homogeneous to ³³² point ($Q_1 \approx 0.2 \text{ Å}^{-1}$), the movement of the molecules ²⁶⁵ heterogeneous crossover [11, 22, 23, 25, 26]. They pro- ³³³ asymptotically obeys Fick's raw ($\tau \propto Q^{-2}$). When Q $_{286}$ posed an anomalous jump diffusion model, where the het- $_{334}$ is higher than Q_1 , the intermediate scattering functions ²⁸⁷ erogeneity is attributed to the distribution of the jump ³³⁵ are well approximated by the KWW function. In addi-288 length. In this interpretation, the heterogeneity is in- 336 tion, the exponent β gradually decreases with increas- $_{289}$ herent in polymers or liquids and not the same as the $_{337}$ ing Q and reaches to 0.5 at the second crossover point ²⁹⁰ dynamical heterogeneity which appears near $T_{\rm g}$. In fact, ³³⁸ $(Q_1 \approx 0.8 \text{ Å}^{-1})$. In this transient region, the relation ²⁹¹ the crossover at Q_2 remains almost unchanged in a wide ³³⁹ $\tau \propto Q^{-2}$ violates and another relation $\tau \propto Q^{-2/\beta}$ is ob-²⁹² temperature range (1.5 $T_{\rm g} < T < 2.5 T_{\rm g}$) [26]. This pic- ³⁴⁰ served in a narrow Q region. Above $Q = Q_2$, τ is again ²⁹³ ture is consistent with our interpretation of Q_2 described ³⁴¹ proportional to Q^{-2} and β remains almost constant at above. 294

Furthermore, some theoretical studies on supercooled 295 ²⁹⁶ liquids have shown that the mean square displacement $_{297}$ $\langle r^2 \rangle$ exhibits a complicated behavior in time; $\langle r^2 \rangle \propto t^2$ at ²⁹⁸ very short times (ballistic motion), $\langle r^2 \rangle \propto t^{\alpha}$ ($\alpha < 1$) at ²⁹⁹ intermediate times (sub-diffusion), and $\langle r^2 \rangle \propto t$ at long ³⁰⁰ times (diffusion). Upon cooling, such a complicated be-301 havior is pronounced and NGP displays non-zero values ³⁰² in the intermediate (sub-diffusive) region [2–4, 6, 8, 9, 11– 13, 16, 17, 25, 26, 30, 33]. In the framework of the mode 303 coupling theory (MCT) [45], the complex behavior in $\langle r^2 \rangle$ 304 is referred to as a crossover from a localized cage motion 305 to diffusion. The results of our experiment and previous 306 works are qualitatively compatible with the MCT pic-307 ture. 308

Finally we comment on the change in β at Q_2 (see 309 ³¹⁰ Fig. 4). The MD simulation works on liquids show that $_{311} \beta$ increases toward 1 when Q is lowered as our experi- $_{312}$ ments, in contrast to polymers in which β remains almost $_{354}$ ³¹³ unchanged. However, their Q-dependency is equivocal. ³⁵⁵ sistance in the SWAN experiment. Travel expenses for ³¹⁴ β changes below $Q \approx 0.5$ Å⁻¹ in glycerol [33], below ³⁵⁶ the experiments at NCNR were supported by General $_{315}Q \approx 0.8$ Å⁻¹ in monomer species of polymer [17], and $_{357}$ User Program for Neutron Scattering Experiments, In-316 gradually changes in a wide Q regime $(Q \le 2 \text{ Å}^{-1})$ in $_{358}$ stitute for Solid State Physics, The University of Tokyo ³¹⁷ water [4] and ortho-terphenyl [9]. While most of experi-³⁵⁹ (proposal No. 15587), at JRR-3, Japan Atomic En-³¹⁸ mental data do not display significant variation of β . It ₃₆₀ ergy Agency, Tokai, Japan. This work utilized facili- $_{319}$ is often difficult to determine β in a wide Q-region due $_{361}$ ties supported in part by the National Science Founda-320 to the limitation of time window in experiments. The 362 tion under Agreement No. DMR-1508249. MN acknowl-321 relation between the non-exponentiality and the Gaus- 363 edges funding support from the cooperative agreement 322 sianity remains an open question and further studies are 364 70NANB15H259 from NIST, U.S. Department of Com-323 required.

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IV. SUMMARY

325 ³²⁶ dependence of self-diffusion process in two types of glass-³⁷¹ available for the purpose.

279 the non-Gaussian parameter (NGP) exhibits a signifi- 327 forming liquids by means of IQENS. Several spectrom-342 0.5.

> Q_1 corresponds to the onset of non-Fickian diffusion 343 344 attributed to dynamical correlation in supercooled liq- $_{345}$ uids. While Q_2 is the crossover point from the Gaus-346 sian to the non-Gaussian character. The present re-³⁴⁷ sults clearly provide the experimental evidence for the ³⁴⁸ two crossovers and Q dependence of β . They are qual-349 itatively consistent with previous MD simulations and 350 IQENS works. We conclude that the two crossovers are ³⁵¹ a generic feature in glass-forming liquids at relatively $_{352}$ higher temperatures than $T_{\rm g}$.

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