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# <sup>1</sup> Combined Experimental and Computational Approach toward the <sup>2</sup> Structural Design of Borosilicate-Based Bioactive Glasses

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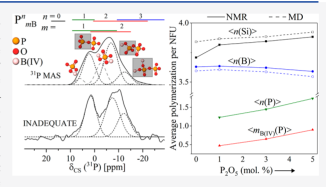
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s **ABSTRACT:** Transitioning beyond a *trial-and-error* based approach 6 for the compositional design of next-generation borosilicate-based 7 bioactive glasses requires a fundamental understanding of the 8 underlying compositional and structural drivers controlling their 9 degradation and ion release *in vitro* and *in vivo*. Accordingly, the 10 present work combines magic-angle spinning (MAS) NMR 11 techniques, MD simulations, and DFT calculations based on 12 GIPAW and PAW algorithms, to build a comprehensive model 13 describing the short-to-medium-range structure of potentially 14 bioactive glasses in the Na<sub>2</sub>O-P<sub>2</sub>O<sub>5</sub>-B<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> system over a 15 broad compositional space. P<sub>2</sub>O<sub>5</sub> preferentially tends to attract 16 network modifier species, thus resulting in a repolymerization of the



17 silicate network and a restructuring of the borate component.  $^{11}B\{^{31}P\}$  and  $^{31}P\{^{11}B\}$  dipolar recoupling experiments suggest that the ability of glasses to incorporate  $P_2O_5$  without phase separation is related to the formation of P-O-B(IV) linkages integrated into the borosilicate glass network. An analogous approach is used for elucidating the local environments of the Na<sup>+</sup> network modifiers. This work, along with future studies aimed at elucidating *composition-structure-solubility/bioactivity* relationships, will lay the foundation for the development of quantitative structure—property relationship (QSPR) models, thus representing a leap forward in the design of functional borosilicate bioactive glasses with controlled ionic release behavior.

#### 1. INTRODUCTION

23 The concept of designing third—generation biomaterials is 24 based on the principle of activating a synchronized sequence of 25 genes at the cellular level by the ionic dissolution products 26 released from the biomaterial during its controlled degradation 27 *in vitro* or *in vivo*. While second-generation biomaterials are 28 designed to be either resorbable or bioactive, these two 29 properties are combined in the third-generation materials, 30 which will, upon implantation, help the body heal itself.<sup>1</sup>

which will, upon implantation, help the body heal itself. While the 45S5 Bioglass composition (24.3Na<sub>2</sub>O–26.9CaO–2.6P<sub>2</sub>O<sub>5</sub>–46.1SiO<sub>2</sub>; mol %), was initially developed as a second-3generation biomaterial, the ability of its ionic dissolution products to stimulate genetic responses in the processes relevant to osteoblast metabolism and bone homeostasis has paved the way for its application as third-generation biomaterial. However, owing to its high tendency toward devitrification (resulting in poor sintering ability—a prerequisite to fabricate 3D porous scaffolds) and slow/poor resorbability, due to the formation of a Si–OH-based passivating gel layer on its surface when in contact with body fluids, its application as a third-2 generation biomaterial has been confined to its use as fine powders/particulates present in toothpaste and dental putty. Therefore, in order to suppress its tendency toward devitrification, and improve its resorbability, B<sub>2</sub>O<sub>3</sub> was introduced into

the silicate network of 45S5 Bioglass, thus resulting in the 46 emergence and development of borosilicate-based bioactive 47 glasses. 11-15

The past 15+ years have experienced a tremendous upsurge in 49 the research interest of boron-containing bioactive glasses as 50 potential candidates for the design and development of novel 51 third-generation biomaterials. Borosilicate glasses, in 52 particular, provide certain advantages over standard silicate-53 based compositions, including (i) a broader glass-forming range, 54 (ii) tunable degradation rates, and (iii) ease of processing into 55 porous three-dimensional scaffolds for application in tissue 56 engineering. Recent studies have further shown their ability 57 to promote angiogenesis and osteogenesis *in vivo*. These 58 features make borosilicate-based bioactive glasses an attractive 59 candidate for application in the treatment of a broad range of 60 skeletal and nonskeletal biomedical problems. However, a major 61 problem that impedes the design of novel borosilicate-based 62

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63 bioactive glasses is our poor understanding of the underlying 64 compositional and structural drivers controlling borosilicate 65 dissolution kinetics when in contact with body fluids. This 66 knowledge gap stems from the fact that the vast majority of 67 research in the field of bioactive glasses have focused on silicate 68 glasses, either based on—or inspired by—45S5 Bioglass.<sup>20–25</sup> 69 Furthermore, as explained in the next section, the principles of 70 silicate glass dissolution/degradation (e.g., composition-struc-71 ture-bioactivity correlations) generally do not apply to borate or 72 borosilicate glasses. As a consequence, the design of borate and 73 borosilicate glasses—especially those that exhibit the desired 74 dissolution behavior—has followed Edisonian approaches 75 involving time-consuming and iterative synthesis-testing cycles. 76 Two notable examples are 13-93B1 and 13-93B3 glasses, derived 77 from a well-known silicate-based bioactive glass (13-93). These 78 compositions have been designed through a trial-and-error 79 approach in the series 6.0Na<sub>2</sub>O-7.9K<sub>2</sub>O-7.7MgO-22.1CaO-80  $1.7P_2O_5-xB_2O_3-(54.6-x)SiO_2$  (mol %), where x = 0 is 13-93 81 and x = 18.2 and 54.6 are 13-93B1 and 13-93B3 compositions, 82 respectively. This partial or complete replacement of SiO<sub>2</sub> with 83 B<sub>2</sub>O<sub>3</sub> was performed with little rationale or revelation of the 84 underlying composition—structure—property relationships. 26,27 85 For a more efficient design of such functional materials, we 86 will require a deeper, more fundamental understanding of the 87 compositional and structural dependence of glass degradation 88 behavior in vitro and in vivo.

# 2. WHY CAN WE NOT APPLY OUR UNDERSTANDING 89 OF THE STRUCTURE OF SILICATE-BASED 90 BIOACTIVE GLASSES TO DESIGN BOROSILICATE 91 GLASSES WITH CONTROLLED ION-RELEASE?

92 The discovery of 45S5 Bioglass was a paradigm shift in the field 93 of biomaterials which later became a benchmark for subsequent 94 research in the field of bioactive glasses. Based on our 95 understanding of the structure and bioactivity of 45S5 Bioglass, 96 the highest bioactivity in silicate-based bioactive glasses arises 97 from a structure dominated by metasilicate chains (Si<sup>2</sup> units), 98 which are occasionally cross-linked through Si<sup>3</sup> units, and are 99 terminated with Si<sup>1</sup> species.<sup>28</sup> Here, and throughout the text,  $T_{mX}^{n}(T, X = Si, B, P)$  refers to the network former unit (NFU) 101 species, where n denotes the number of bridging oxygen (BO) 102 atoms and  $m \le n$  specifies the number of linkages to other NFUs 103 of type X. It should be noted here that when correlating the 104 structural makeup of silicate glasses with their bioactivity, the 105 latter was previously considered as the rate of formation of 106 hydroxyapatite on the glass surface when in contact with body 107 fluids.<sup>21</sup> However, hydroxyapatite formation is no longer 108 considered to be an initial marker of bioactivity and the ability 109 to achieve a tunable degradation rate with controlled ion-release 110 is instead a prerequisite for designing third-generation bioactive 111 glasses suitable for tissue engineering. Accordingly, silicate 112 glasses with highly polymerized glass structures, for example, 113 glasses consisting of high fractions of Si<sup>3</sup> or Si<sup>4</sup> units, may exhibit 114 well-controlled ionic release properties while at the same time 115 having lower bioactivity (i.e., glass 55S) or even becoming 116 bioinactive (i.e., glass 65S).<sup>2</sup>

Furthermore,  $P_2O_5$ , though not strictly necessary for list bioactivity, plays a vital role in silicate glasses to enhance their bioactivity when present in small concentrations (for example, 2.5 mol % in 45S5 Bioglass). On a structural level, the enhanced bioactivity of  $P_2O_5$ -containing silicate glasses has been attributed to the presence of loosely bound orthophosphate

units  $(P^0)$  in the glass structure whose relatively fast initial 123 release from the glass into the solution enhances the bone- 124 bonding ability of the glass. 125

When compared with the structure of silicate-based bioactive 126 glasses, the structure of borosilicate-based glasses is much more 127 complex.<sup>31,32</sup> Similar to silicate-based glass chemistries, the 128 dissolution behavior of borosilicate-based bioactive glasses is 129 controlled by the short-to-medium-range order in the glass 130 structure. As B<sub>2</sub>O<sub>3</sub> enters the silica glass network, in addition to 131 the introduction of new short-range order species (i.e., three- 132 and four- coordinate boron species), the intermediate-range 133 associations between NFUs in borosilicate glasses (e.g., Si<sup>4</sup>-O- 134  $B^4$ ,  $Si^4-O-B^3$ ,  $B^3-O-B^4$ , etc. linkages) further complicate the 135 glass structure. 33-36 These associations are expected to have a 136 significant impact on the dissolution kinetics and bioactivity of 137 these glasses. For instance, common boron NFU linkages in 138 borosilicate glasses have lower enthalpies of hydrolysis, i.e., Si<sup>4</sup> – 139  $O-B^3$  (-3.03 kJ/mol) and  $B^3-O-B^3$  (-16.98 kJ/mol), as 140 compared to pure silica linkages (Si<sup>4</sup>–O–Si<sup>4</sup> linkages: 16.61 kJ/ 141 mol),<sup>37</sup> resulting in the greater degradation rates typically 142 observed upon B<sub>2</sub>O<sub>3</sub> substitution into silicate-based bioactive 143 glasses. <sup>17</sup> The addition of P<sub>2</sub>O<sub>5</sub> further adds to the composi- 144 tional and structural complexity of this glass system by forming 145 B-O-P linkages. 38,39 The previously reported impacts of P<sub>2</sub>O<sub>5</sub> 146 additions to borosilicate glasses include an increased degree of 147 polymerization of the silica and phosphate species and P–O–B  $_{148}$  intermixing in the network.  $^{39-41}$   $P_2O_5$  is generally known as a  $_{149}$ cation scavenger in mixed network former glasses, attracting 150 large amounts of modifier via multiple charged orthophosphate 151 and diphosphate anions.<sup>38-40</sup> However, the impact of these 152 structural units and linkages on the dissolution kinetics of glasses 153 is still largely unknown. Thus, it is difficult, if not impossible, to 154 design next-generation borosilicate-based bioactive glasses using 155 an approach based on the structure of silicate glasses.

In the context of the above-discussed rationale, the present 157 contribution aims to combine the strengths of both experimental 158 and computational materials science to uncover the short-to- 159 medium-range structure of potentially bioactive glasses in the 160 system  $Na_2O-P_2O_5-B_2O_3-SiO_2$  system over a broad composi- 161 tional space with distinct structural features. Although we 162 understand that most bioactive glass compositions contain CaO 163 as a major component, we have selected a calcium-free glass 164 system for the following reasons:

- (i) Competition between  $Na^+$  and  $Ca^{2+}$  (i.e., in a  $Na_2O$  166  $CaO-P_2O_5-B_2O_3-SiO_2$  system) nonframework cations 167 for association with NFU units (B, P, Si) would add 168 significant additional complexity while interpreting glass 169 structural speciation.
- (ii) The structural role of Ca<sup>2+</sup> in these glasses would be 171 challenging to investigate in the absence of prior in-depth 172 knowledge of the CaO-free system, given the extreme 173 difficulty in measuring <sup>43</sup>Ca NMR spectra in glasses 174 without an isotopic enrichment and/or an elevated Ca- 175 presence in the sample. <sup>42-44</sup> Additionally, the tendency of 176 glasses containing both sodium and calcium to decrease 177 the resolution of <sup>29</sup>Si and <sup>31</sup>P MAS NMR spectra further 178 complicates their interpretation due to the creation of 179 multiple chemical environments. <sup>28</sup>
- (iii) The glass-forming ability of compositions containing both  $^{181}$  CaO and  $P_2O_5$  is anticipated to be highly limited, due to  $^{182}$  the high cation field strength of  $Ca^{2+}$  and the high  $^{183}$

Table 1. All Batched Compositions of Studied Glasses Compared to Selected Experimental Compositions (in Brackets), as Analyzed via ICP-OES ( $\pm 0.5$  mol. %), with Density ( $\rho$ ) and Molar Volume ( $V_{\rm M}$ ) Also Displayed

		ba				
sample ID	Na <sub>2</sub> O	$P_2O_5$	$B_2O_3$	SiO <sub>2</sub>	$\rho$ (g/cm <sup>3</sup> ) ( $\pm$ 0.3%)	$V_{\mathrm{M}}$ (±0.3%) (cm <sup>3</sup> /mol
PB0	25.0 [25.3]	-	30.0 [29.7]	45.0 [45.0]	2.519	25.18
PB1-P1	24.6	1.0	29.5	45.0	2.509	25.58
PB1-P3	23.6	3.0	28.4	45.0	2.479	26.50
PB1-P5	22.7	5.0	27.27	45.0	2.462	27.30
PB2-P1	24.8	1.0	29.7	44.6	2.518	25.50
PB2-P3	24.3 [25.9]	3.0 [3.0]	29.1 [29.0]	43.7 [42.2]	2.507	26.24
PB2-P5	23.8 [25.8]	5.0 [5.0]	28.5 [29.1]	42.8 [40.0]	2.476	27.20
PB3-P1	25.0	1.0	30.0	44.0	2.510	25.59
PB3-P4	25.0	4.0	30.0	41.0	2.485	26.84
PB3-P7	25.0	7.0	30.0	38.0	2.456	28.15
MB0	25.0 [25.1]	_	25.0 [25.0]	50.0 [49.9]	2.531	24.87
MB1-P1	24.5	1.0	24.5	50.0	2.519	25.29
MB1-P3	23.5	3.0	23.5	50.0	2.504	26.05
MB1-P4	23.0	4.0	23.0	50.0	2.490	26.50
MB2-P1	24.8	1.0	24.8	49.5	2.533	25.16
MB2-P3	24.3 [26.0]	3.0 [3.0]	24.3 [23.9]	48.5 [47.1]	2.518	25.94
MB2-P5	23.8 [24.7]	5.0 [5.0]	23.8 [23.9]	47.5 [46.3]	2.488	26.88
MB3-P1	25.0	1.0	25.0	49.0	2.519	25.31
MB3-P3	25.0	3.0	25.0	47.0	2.510	26.05
MB3-P5	25.0	5.0	25.0	45.0	2.487	26.95
PA0	25.0 [25.1]	_	20.0 [20.1]	55.0 [54.8]	2.527	24.72
PA1-P1	24.4	1.0	19.6	55.0	2.501	25.28
PA1-P3	23.3	3.0	18.7	55.0	2.497	25.94
PA2-P1	24.8	1.0	19.8	54.5	2.526	25.04
PA2-P3	24.3 [25.1]	3.0 [3.1]	19.4 [19.3]	53.4 [52.6]	2.513	25.80
PA3-P1	25.0	1.0	20.0	49.0	2.515	25.16
PA3-P3	25.0	3.0	20.0	47.0	2.506	25.91

tendency of P<sub>2</sub>O<sub>5</sub> to phase separate/crystallize when introduced into silica-based glasses. <sup>39,40,45</sup>

- (iv) Although CaO is an important component in bioactive glasses and bioceramics for promoting intracellular and extracellular bodily responses, <sup>46</sup> CaO-free bioactive glasses and bioceramics displaying bioactivity *in vitro* and *in vivo* have also been proposed in the literature. <sup>47–54</sup>
- (v) The inclusion of CaO into the glass system would significantly increase the number of experiments. It would be difficult to comprehend all the results and their analysis in one article.

The overarching goal of the study is to develop a comprehensive model elucidating the structure of sodium phospho-borosilicate glasses. Our future studies will focus on understanding the impact of nonframework cation mixing, with special emphasis on elucidating the impact of Ca<sup>2+</sup> on (i) the short-to-medium-range ordering in the glass structure, and (ii) the kinetics and mechanisms of chemical degradation and ion release from these glasses. The objective of the current and forthcoming articles on this topic is to lay the foundation for a rational design of borosilicate-based bioactive glasses based on the quantitative structure—property relationships (QSPR) approach.

# 3. EXPERIMENTAL SECTION

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3.1. Glass Composition Design. In order to design glasses over a broad compositional space with a variety of structural features, the baseline glasses were selected with respect to their  $Na_2O/B_2O_3$  (=R) ratio. This ratio has been shown to have a significant impact on the short-to-medium-range ordering in the

structure of borosilicate glasses as has been shown by Martens 212 and Muller-Warmuth, and Du and Stebbins. Accordingly, 213 three baseline glasses with compositions (mol %) (i) 25 Na<sub>2</sub>O- 214 30 B<sub>2</sub>O<sub>3</sub>-45 SiO<sub>2</sub> (R=0.83), (ii) 25 Na<sub>2</sub>O-25 B<sub>2</sub>O<sub>3</sub>-50 SiO<sub>2</sub> 215 (R=1), and (iii) 25 Na<sub>2</sub>O-20 B<sub>2</sub>O<sub>3</sub>-55 SiO<sub>2</sub> (R=1.25) were 216 selected in the perboric (PB, R<1), metaboric (MB, R=1), and 217 peralkaline (PA, R>1) homogeneous glass-forming regions of 218 the sodium borosilicate ternary diagram.

Further, to unambiguously understand the impact of  $P_2O_5$  on 220 the structure of sodium borosilicate glasses, the former was 221 introduced in the baseline glass systems using three different 222 approaches, with the value of R being constant. The substitution 223 schemes (Schemes 1–3) for metaboric (MB) glasses serve as 224 representative examples for all the glasses investigated in the 225 present study.

series MB1: 
$$(25 - x/2)$$
Na<sub>2</sub>O $-x$ P<sub>2</sub>O<sub>5</sub> $-(25 - x/2)$ B<sub>2</sub>O<sub>3</sub> $-50$ SiO<sub>2</sub>

series MB2: 
$$xP_2O_5-(100-x)(25Na_2O-25B_2O_3-50SiO_2)$$

series MB3: 
$$25\text{Na}_2\text{O} - xP_2\text{O}_5 - 25\text{B}_2\text{O}_3 - (50 - x)\text{SiO}_2$$

Accordingly, each borosilicate glass system, i.e., perboric (PB),  $^{230}$  metaboric (MB), and peralkaline (PA), has been further divided  $^{231}$  into three series of glasses based on the scheme of  $^{20}$ 5  $^{232}$ 

233 substitution and have been labeled in accordance with the above. 234 The batched  $P_2O_5$  contents (x) in the investigated glasses vary 235 between 0 and 9 mol %. However, only the batched 236 compositions resulting in visibly transparent glasses were 237 structurally investigated.

**3.2. Glass Synthesis.** High purity powders of SiO<sub>2</sub> (Alfa 239 Aesar; 99.5%),  $H_3BO_3$  (Alfa Aesar;  $\geq$  98%),  $Na_2SiO_3$  (Alfa 240 Aesar; >99%), and Na<sub>2</sub>HPO<sub>4</sub> (Fisher Chemical; >99%) were 241 used as precursors. Batches corresponding to 70 g of oxides were 242 melted in Pt-Rh crucibles for 1 h in air at temperatures between 243 1400 and 1500 °C and quenched on a metallic plate. The 244 amorphous nature of glasses was confirmed by X-ray diffraction (XRD) (PANalytical-X'Pert Pro; Cu K $\alpha$  radiation;  $2\theta$  range  $246\ 10-90^{\circ}$ ; step size  $0.01313^{\circ} \, s^{-1}$ ). The glasses were then annealed 247 for 1 h near their glass transition temperatures ( $T_{\rm g}^*$  – 50 °C, 248 where  $T_{\rm g}^*$  is predicted from the SciGlass database <sup>56</sup>) and slowly 249 cooled to room temperature (see ref 18 for details). Residual 250 stresses of less than 10 MPa were measured using a polariscope. The experimental compositions of the selected glasses 252 (especially those with high Na<sub>2</sub>O, B<sub>2</sub>O<sub>3</sub>, and P<sub>2</sub>O<sub>5</sub> concen-253 tration) were analyzed using inductively coupled plasma-254 optical emission spectroscopy (ICP-OES; PerkinElmer Opti-255 ma 7300 V) for B<sub>2</sub>O<sub>3</sub>, P<sub>2</sub>O<sub>5</sub>, and SiO<sub>2</sub>, and flame emission 256 spectroscopy for Na<sub>2</sub>O (PerkinElmer Flame Emission Analyst 257 200). Table 1 presents the compositions of all the glasses 258 investigated in the present study. The density of glasses was 259 measured using Archimedes' method for at least three 260 specimens per glass composition weighed in both air and d-261 limonene.

3.3. Structural Analysis by NMR Spectroscopy. NMR spectra were measured on (i) an Agilent 240-MR DD2 spectrometer (5.7 T) using 3.2, 4.0, and 7.5 mm MAS NMR probes, (ii) Bruker Avance Neo 500 and 600 MHz spectrometers (11.7 and 14.1 T, respectively) using 2.5 and 4.0 mm MAS NMR probes, (iii) an Agilent DD2 system at 16.4 T equipped with a 3.2 mm MAS NMR probe, and (iv) an Agilent VNMRs system at 11.7 T equipped with a 5 mm MAS NMR probe. Table S1 specifies the conditions used for the various single resonance measurements.

One-dimensional refocused INADEQUATE experiments were employed on selected samples to probe the connectivity between phosphate units. This technique uses double-quantum filtering, based on homonuclear indirect spin—spin ("J-coupling"), for selective detection of P–O–P linked Avance Neo 600 MHz spectrometer at 14.1 T (243.0 MHz), and a Bruker DSX 500 console interfaced with a 4.7 T magnet, with a 280 2.5 (4.0) mm probe using a spinning frequency of 15.0 (12.0) kHz,  $\pi/2$  pulse lengths near 1.6 (4.0)  $\mu$ s, recycle delays of 5–10 s, and signal averaging over at least 4000 (24000) acquisitions. The double quantum coherence buildup time was set to 16 ms, corresponding to a  ${}^2J({}^{31}P-{}^{31}P)$  coupling constant of 30 Hz.

All the single resonance MAS NMR spectra were analyzed using the DMFit software, <sup>59</sup> utilizing the CzSimple model for <sup>23</sup>Na MAS NMR spectra, Gauss/Lorentz functions for <sup>31</sup>P and <sup>288</sup> <sup>29</sup>Si MAS NMR spectra, and the "Q MAS 1/2" model and Gauss/Lorentz functions, respectively, for the <sup>11</sup>B resonances of <sup>290</sup> 3- and 4-coordinated boron species.  $N_4$  values were determined <sup>291</sup> from the fractional areas of the peaks attributed to the four-<sup>292</sup> coordinated boron species, with a small correction for the <sup>293</sup> overlapping satellite transition of the B(IV) species. <sup>60</sup>

Dipolar interactions between  $^{11}$ B,  $^{31}$ P, and  $^{23}$ Na nuclei were probed using  $^{11}$ B $\{^{31}$ P $\}$ ,  $^{23}$ Na $\{^{31}$ P $\}$ , and  $^{31}$ P $\{^{23}$ Na $\}$  rotational

echo double resonance (REDOR) experiments, in addition to 296 <sup>31</sup>P{<sup>11</sup>B} rotational echo adiabatic passage double resonance <sub>297</sub> (REAPDOR) spectroscopy, on selected samples. All REDOR 298 experiments involving <sup>23</sup>Na-<sup>31</sup>P nuclear interactions were 299 conducted on a Bruker DSX 400 spectrometer using a 4.0 mm 300 probe at 9.4 T. <sup>11</sup>B-<sup>31</sup>P double resonance measurements were 301 conducted on a Bruker Avance Neo 600 MHz spectrometer 302 using a 2.5 mm probe at 14.1 T. Table S1 specifies the conditions 303 used for REDOR and REAPDOR experiments. The normalized 304 REDOR and REAPDOR signal intensities  $\Delta S = (S_0 - S)/S_0$  305 (where S and  $S_0$  are the signals (i) with and (ii) without 306 recoupling  $\pi$  pulses, respectively) was plotted as a function of 307 dipolar mixing time  $(NT_r)$ , where N is the number of rotor 308 cycles and  $T_r$  is the rotor period. REDOR experiments were 309 performed using a rotor-synchronized spin echo sequence using 310  $\pi$  pulses as shown in Table S1. The compensation pulse scheme 311 was used,  $^{61}$  and  $\pi$  pulses on the  $^{31}$ P channel were phase cycled  $_{312}$  according to the XY-4 scheme.  $^{62}$  Following previously  $_{313}$  established procedures,  $^{61,63}$  dipolar second moments  $(M_{2(S-I)})$   $_{314}$ (where S represents the observed nucleus and I represents the 315 nonobserved nucleus) were determined by fitting the initial part 316 of the REDOR curves ( $\Delta S/S_0 \leq 0.20$ ) using the parabolic 317 approximation shown in eq 1:

$$\frac{\Delta S}{S_0} = \frac{4}{3\pi^2} M_{2(S-I)} (NT_r)^2 \tag{1)}{}_{319}$$

The final  $M_{2(S-I)}$  values are obtained by calibration with 320 experimental data on the crystalline model compounds BPO<sub>4</sub> 321 and Na<sub>3</sub>P<sub>3</sub>O<sub>9</sub>, for which the theoretical second moments can be 322 calculated from crystallographic information. 64,65 The REAP- 323 DOR pulse sequence is typically used in the  $S\{I\}$  case where the 324 I-nuclei are quadrupolar, such as <sup>11</sup>B, producing more efficient 325 dephasing by applying an adiabatic passage pulse lasting one- 326 third of a rotor period in the middle of the dipolar recoupling 327 period.<sup>66</sup> As the signal-to-noise ratio of these experiments is 328 limited by the long-spin-lattice relaxation times of the <sup>31</sup>P- <sup>329</sup> observe nuclei, data were measured for two or three dipolar 330 mixing times only, using rotor-synchronized  $\pi$  pulses in the <sup>31</sup>P <sub>331</sub> channel and an adiabatic passage pulse on the <sup>11</sup>B channel. 332 Additionally, a saturation comb of 60 90° pulses ensured a 333 stationary initial magnetization at the beginning of each 334 experiment. Simulated REAPDOR curves for each sample 335 were generated using the SIMPSON program package, 67 taking 336 into account the experimentally determined spin-spin inter- 337 action parameters and the experimental conditions.

3.4. Molecular Dynamics Simulations. Classical molec- 339 ular dynamics simulations have been employed to obtain further 340 insight into the structure of the glass series PB2, MB2, and PA2, 341 and on the glass PB3-P7, using the batched compositions as 342 reported in Table 1. Models containing about 3500 atoms have 343 been generated (three replicas for each composition) by using 344 the melt-quench approach.<sup>68</sup> The exact number of atoms and 345 box dimensions are reported in Table S2 for these compositions, 346 as well as for the compositions (mol %): (i) 55Na<sub>2</sub>O-45P<sub>2</sub>O<sub>5</sub> 347 and (ii)  $40\text{Na}_2\text{O} - 18\text{B}_2\text{O}_3 - 42\text{P}_2\text{O}_5$ , which have been similarly 348 modeled for the NMR calculations (described below). The shell 349 model force-fields have been used to describe the interatomic 350 interactions between ionic pairs. In this model, which has been 351 demonstrated to reproduce better the medium-range structure 352 of oxide glasses (especially in terms of  $T^n$  distributions and 353 intertetrahedral bond angle distributions), 69-71 the more 354 polarizable ions (oxygen in this case) are represented by a 355

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ass6 massive core connected to a massless shell by a harmonic spring. A charge is assigned to both the core and shell. The functional ss form presented in eq 2, which has been already used in previous studies, that also been utilized in the present study.

$$U(r_{ij}, r_{c-s}, \theta_{ijk}) = k \frac{q_i q_j}{r_{ij}^2} + A_{ij} e^{-(r_{ij}/\rho_{ij})} - \frac{C_{ij}}{r_{ij}^6} + \frac{1}{2} k_s (r_{core-shell})^2 + \frac{1}{2} k_b (\theta_{ijk} - \theta_{ijk}^0)^2 \exp\left(-\frac{r_{ij}}{\rho} - \frac{r_{jk}}{\rho}\right)$$
(2)

361 Here the first term describes Coulombic interactions between all 362 ions (the core and shell belonging to the same ion are filtered 363 out), the second term is a Buckingham function applied between 364 the cation cores and oxygen shells, and the third term represents 365 the harmonic spring connecting the core and shell of the same 366 oxygen ions whereas the last term is a three-body interaction 367 used to constrain the O–Si–O and O–P–O angles to 109°.

All the parameters used are included in Table S3 of the 369 Supporting Information. It is important to highlight that all the 370 parameters have been already used in previous investigations 371 with the exception of the B-O parameter sets. 20,69,73 The latter 372 have been refined starting from those developed by Edén et al. 41 373 to be consistent with the other parameters and to better 374 reproduce the  $N_4$  fraction in borosilicate glasses with  $[B_2O_3]/$  $375 [SiO_2] \le 0.33$ . The leapfrog algorithm encoded in the 376 DL\_POLY2.14 package<sup>74</sup> has been used to integrate the 377 equations of motion with a time step of 0.2 fs. The initial 378 configurations were generated by randomly placing the number 379 of atoms in a cubic box, whose dimensions were constrained by 380 the experimental densities. The systems were heated and held at 381 3200 K for 100 ps in the NVT ensemble ensuring a suitable 382 melting of the samples. The liquids were then cooled to 300 K at 383 a nominal cooling rate of 5 K/ps. The resulting glass structures 384 were subjected to a final equilibration run of 200 ps. Velocity 385 scaling was applied at every step during the quenching of the 386 melt to control the kinetic energy (temperature) of the shells. 387 Coulomb interactions were calculated by the Ewald summation 388 method with a cutoff of 8 Å, whereas short-range cutoff values of 389 7.5 Å were used for the (vdW) short-range interactions.

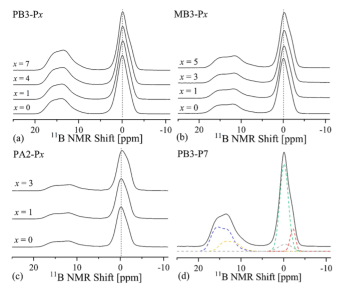
**3.5. NMR Parameter Calculations.** To further guide the 391 interpretation of the NMR experiments, we have also computed 392 the NMR parameters of <sup>17</sup>O, <sup>31</sup>P, <sup>29</sup>Si, <sup>11</sup>B, and <sup>23</sup>Na nuclei for 393 the PB2-P5,  $55Na_2O-45P_2O_5$ , and  $40Na_2O-18$  B<sub>2</sub>O<sub>3</sub> $-42P_2O_5$ 394 glass compositions. Three models of this glass containing 374 395 atoms each have been generated as was described above. 396 Magnetic shielding and EFG tensors of the various NFUs 397 present were computed with the NMR-CASTEP<sup>75</sup> density 398 functional theory (DFT) code using the GIPAW<sup>76</sup> and PAW<sup>7</sup> 399 algorithms, respectively. The generalized gradient approxima-400 tion (GGA) PBE<sup>78</sup> functional was employed, and the core-401 valence interactions were described by ultrasoft pseudopoten-402 tials generated on the fly. For <sup>17</sup>O, the 2s and 2p orbitals were 403 considered as valence states with a core radius of 1.3 Bohr; for 404 <sup>29</sup>Si and <sup>31</sup>P, a core radius of 1.8 Bohr was used with 3s and 3p 405 valence orbitals; for <sup>23</sup>Na, a core radius of 1.3 Bohr was used with 406 2s, 2p, and 3s valence orbitals, while for 11B, a core radius of 407 1.405 Bohr was used with 2s and 2p valence states. For the PAW 408 and GIPAW calculations we used two projectors in each s and p 409 angular momentum channel for O and B, and in each s, p, and d 410 channel for Si and Na. Before computing the NMR parameters,

#### 4. RESULTS AND DISCUSSION

analyzed using the SoSNMR software.83

4.1. Glass Formation and Bulk Properties. Among the 428 samples within a range of x = 0 to 9 mol % for each series, those 429 ultimately selected for study passed the criteria of being 430 transparent in appearance after annealing and showing an 431 amorphous character in XRD (Figure S1). Perboric (Na/B < 1) 432 glasses can incorporate  $P_2O_5$  in amounts up to x = 5-7 mol % 433 while in metaboric and peralkaline glasses the limits are x = 4-5 434 mol % and x = 3 mol %, respectively. Peralkaline glasses 435 synthesized with more than 3 mol % P2O5 exhibit evidence of 436 crystalline Na<sub>4</sub>P<sub>2</sub>O<sub>7</sub> phase formation in XRD—see Figure S1— 437 in addition to being visibly phase-separated/crystallized. ICP- 438 OES (see Table 1) analyses conducted on representative 439 samples show a close agreement of their B<sub>2</sub>O<sub>3</sub> and P<sub>2</sub>O<sub>5</sub> contents 440 with their batched compositions (within ±0.6 mol %). The 441 analyzed concentrations of Na2O and SiO2 in the baseline 442 glasses are also close to their batched values (within  $\pm 0.5$  mol 443 %). However, larger variations in the values of Na<sub>2</sub>O and SiO<sub>2</sub> 444 can be seen in  $P_2O_5$ -containing glass samples (x = 3-5 mol %), 445 with differences ranging between ±0.7 and 2.7 mol %. The 446 experimental densities and molar volumes of the synthesized 447 glasses are also presented in Table 1. For the baseline glasses, 448 density values agree with previous trends,84 while upon 449 successive introduction of P2O5, the molar volume tends to 450 increase, reflecting the larger size of the phosphate component. 451

4.2. 11B MAS NMR and 11B{31P} REDOR. Figure 1a-c 452 ft presents the <sup>11</sup>B MAS NMR spectra of glasses in the series PB3, 453 MB3, and PA2 as representatives for all the glasses investigated 454 in the present study. The <sup>11</sup>B MAS NMR spectra of all the other 455 glasses have been presented in Figure S2. All the spectra show a 456 structured line shape near 14 ppm, reflecting strong quadrupolar 457 perturbations on the <sup>11</sup>B signals of three-coordinated boron, and 458 sharper features near 0 ppm, arising from four-coordinated 459 boron, B(IV), for which quadrupolar interactions are signifi- 460 cantly weaker. To obtain satisfactory fits one must assume at 461 least two B(III) and two B(IV) components each (e.g., see 462 Figure 1d and Table S4). While this deconvolution should be 463 considered artificial, it serves well for extracting reliable N<sub>4</sub> 464 values from these spectra. Regarding assignments, we expect 465 B<sup>3</sup>(III) with three bridging oxygen species within ring and 466 nonring units, but also anionic B<sup>2</sup>(III) units featuring two 467 bridging and one nonbridging oxygen atoms. For the P2O5-free 468 baseline glasses, PB0, MB0, and PA0, the fraction of B(IV) units, 469  $N_4$ , agrees very well with the previous results on borosilicate 470 glasses, and predictions based on R and the  $SiO_2/B_2O_3$  (=K) 471



**Figure 1.**  $^{11}$ B MAS NMR spectra of series (a) PB3, (b) MB3, and (c) PA2 glasses. Series PB3 spectra were collected at 16.4 T while series MB3 and PA2 spectra were collected at 14.1 T. (d) Exemplary line shape deconvolution for the central transition of the  $^{11}$ B MAS NMR spectra of the PB3-P7 glass. Each  $^{11}$ B MAS NMR spectrum was fitted with two Q MAS 1/2 components for the B(III) resonances and three Gauss/Lorentz functions for the B(IV) resonances. The minor fitted peak displayed near 0 ppm represents the central peak of the satellite transition manifold of the B(IV) resonances, which overlaps with the MAS peaks of the central transition, and whose area needs to be considered when extracting  $N_4$  values from these spectra.

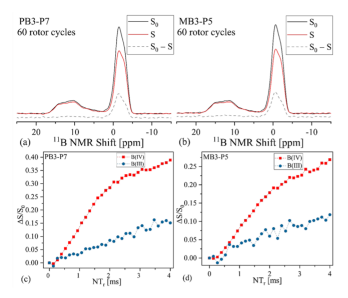
472 ratio.<sup>34</sup> In PB and MB glasses, increasing phosphate content 473 results in a moderate decrease in the fraction of four-coordinated 474 boron,  $N_4$ , which can be attributed to the need for extra charge 475 compensation required for the anionic phosphate species 476 formed. In the case of the PA glasses where significant amounts 477 of  $B^2(III)$  units are expected, the observed invariance of  $N_4$  may 478 be due to the depletion of both anionic four-coordinated B(IV) 479 and anionic three-coordinated B<sup>2</sup>(III) units. The chemical shifts 480 of the  $B^4(IV)$  units, near 0 ppm and -2.5 ppm, are close to those previously reported for  $B^4_{3Si,1B}$  and  $B^4_{4Si}$  units, respectively. 39,85 482 While the chemical shift near -2.5 ppm may have come from an 483 increased number of Si neighbors around borate units for glasses 484 with x = 0, we can alternatively explain it to signify the formation 485 of some B-O-P linkages, which enhances the upfield shoulder 486 somewhat in spectra of glasses containing higher amounts of 487 P<sub>2</sub>O<sub>5</sub>. This question has been explored further using chemical 488 shift calculations, MD simulations, and REDOR experiments, as 489 discussed below. Computed isotropic chemical shifts  $\delta_{ ext{CS}}^{ ext{iso}}$  of all 490 B<sup>3</sup>(III) and B<sup>4</sup>(IV) species range between 14 and 18 ppm and 491 between -2.3 and +1.1 ppm, respectively, in excellent 492 agreement with the experimental values. Table 2 reveals that 493 the  $\delta_{CS}^{iso}$  values of both  $B^3(III)$  and  $B^4(IV)$  species are expected 494 to decrease when B(IV) units in the second coordination 495 spheres are replaced by B(III), Si, and/or P. It is also worth 496 noting that in our models, the most negative isotropic chemical 497 shifts of B<sup>4</sup>(IV) are found for species surrounded by either one 498 B(IV), one B(III), and two P atoms (-1.5 ppm), or by four Si 499 NFUs (-2.3 ppm). Thus, on the basis of these calculations, the so chemical shift near -2.5 ppm can be ascribed to the formation of 501 either multiple B(IV)-O-Si or of multiple B(IV)-O-P 502 linkages or a combination of both.

Table 2. Calculated Isotropic Chemical Shifts of Boron, Silicon, and Phosphorus NFUs Found in the PB2-P5 Glass Modeled at the MD-GIPAW Level

	a ira ( )	2 (2 )
species	$\delta_{ ext{CS}}^{ ext{iso}}\left( ext{ppm} ight)$	$C_{\mathbb{Q}}(MHz)$
$P^1(B(IV))$	$8.2 \pm 4.8$	_
$P^1(B(III))$	$7.1 \pm 5.1$	-
$P^1(Si)$	$4.3 \pm 1.8$	-
$P^1(P)$	$3.8 \pm 3.0^{a}$	-
$P^2(2B(IV))$	$-3.5 \pm 2.6$	_
$P^2(2Si)$	$-10.8 \pm 3.7$	_
$P^2(SiB(IV))$	$-11.5 \pm 1.7$	_
$P^2(B(IV)P)$	$-14.8 \pm 4.4^{b}$	_
$P^3(3B(IV))$	$-18.8 \pm 3.0$	_
$P^3(Si2B(IV))$	$-23.2 \pm 2.0$	_
$Si^4(4B(IV))$	$-82.0 \pm 4.1$	_
$Si^4(3B(IV)B(III))$	$-92.0 \pm 4.0$	_
$Si^4(SiB(III)2B(IV))$	$-95.5 \pm 2.0$	_
$Si^4(2Si2B(IV))$	$-95.9 \pm 1.0$	_
$Si^4(2B(IV)2B(III))$	$-98.6 \pm 3.0$	_
$Si^4(B(III)B(IV)SiP)$	$-99.6 \pm 2.5$	_
$Si^4(Si3B(IV))$	$-100.2 \pm 4.0$	_
$Si^4(B(IV)3B(III))$	$-100.3 \pm 3.5$	_
$Si^4(3SiB(IV))$	$-101.1 \pm 3.0$	_
$Si^4(2SiB(IV)B(III))$	$-101.6 \pm 1.1$	_
$Si^4(Si2B(III)B(IV))$	$-101.6 \pm 3.0$	_
$Si^4(B(IV)2SiP)$	$-102.6 \pm 2.0$	_
Si <sup>4</sup> (2Si2B(III))	$-103.2 \pm 1.0$	_
Si <sup>4</sup> (4Si)	$-107.2 \pm 4.0$	_
Si <sup>4</sup> (4B(III))	$-107.6 \pm 2.0^{c}$	_
Si <sup>4</sup> (3SiB(III))	$-109.2 \pm 2.0$	_
$B^3(2B(III)Si)$	$17.8 \pm 0.5$	$2.85 \pm 0.1$
$B^3(3B(IV))$	$17.2 \pm 2.0$	$2.71 \pm 0.1$
$B^3(B(IV)B(III)Si)$	$16.9 \pm 1.8$	$2.67 \pm 0.1$
$B^3(2B(IV)Si)$	$16.6 \pm 1.3$	$2.65 \pm 0.1$
$B^3(2B(IV)B(III))$	$16.1 \pm 1.0$	$2.68 \pm 0.1$
$B^3(2B(IV)P)$	$15.9 \pm 1.4$	$2.73 \pm 0.1$
$B^3(B(IV)2Si)$	$15.1 \pm 1.0$	$2.67 \pm 0.1$
$B^3(B(III)2Si)$	$14.6 \pm 1.6$	$2.71 \pm 0.1$
$B^4(3B(IV)B(III))$	$1.1 \pm 0.5$	$0.26 \pm 0.05$
$B^4(B(IV)3B(III))$	$0.6 \pm 0.4$	$0.44 \pm 0.05$
$B^4(2B(III)B(IV)Si)$	$0.5 \pm 0.4$	$0.60 \pm 0.05$
$B^4(3B(IV)Si)$	$-0.2 \pm 0.3$	$0.37 \pm 0.05$
$B^4(2B(IV)B(III)Si)$	$-0.3 \pm 0.3$	$0.41 \pm 0.05$
$B^4(4B(III))$	$-0.8 \pm 0.4$	$0.19 \pm 0.05$
$B^4(B(IV)3Si)$	$-0.8 \pm 0.2$	$0.35 \pm 0.05$
$B^4(3B(III)Si)$	$-0.9 \pm 0.3$	$0.17 \pm 0.05$
$B^4(B(III)3Si)$	$-1.0 \pm 0.3$	$0.38 \pm 0.05$
$B^4(B^4B(III)2P)$	$-1.5 \pm 0.4$	$0.20 \pm 0.05$
$B^4(4Si)$	$-2.3 \pm 0.4$	$0.36 \pm 0.05$

 $^a$  As calculated from MD simulations of a 55Na<sub>2</sub>O-45P<sub>2</sub>O<sub>5</sub> glass  $^b$  As calculated from MD simulations of a 40Na<sub>2</sub>O-18 B<sub>2</sub>O<sub>3</sub>-42P<sub>2</sub>O<sub>5</sub> glass  $^c$  As calculated from MD simulations of a 25Na<sub>2</sub>O-56.25B<sub>2</sub>O<sub>3</sub>-18.75SiO<sub>2</sub> glass  $^{71}$ 

In order to quantify the extent of B-O-P linkages,  $^{11}B(^{31}P)$  503 REDOR experiments, which probe  $^{11}B-^{31}P$  magnetic dipole—504 dipole interactions, have been performed on selected samples. 505 Parts a and b of Figure 2 compare the Fourier transformed  $^{11}B$  506 f2 MAS NMR spectra in the absence ( $S_0$ ) and presence (S) of 507 dipolar recoupling and the corresponding difference spectra ( $S_0$  508 - S) for the samples PB3-P7 and MB3-P5, respectively. Such 509 REDOR experiments are instrumental for peak identification 510



**Figure 2.** Fourier transforms of the  $^{11}\text{B}$  MAS spin echoes obtained in a  $^{11}\text{B}\{^{31}\text{P}\}$  REDOR experiment after 60 rotor cycles ( $NT_r=4.0$  ms): reference signal,  $S_0$ , signal with dipolar dephasing,  $S_r$ , and REDOR difference spectrum  $S_0-S_r$ , for (a) PB3-P7 glass and (b) MB3-P5 glass. (c and d)  $^{11}\text{B}\{^{31}\text{P}\}$  REDOR curves ( $\Delta S/S_0$  vs  $NT_r$ ) for PB3-P7 and MB3-P5 samples, respectively. Red squares and blue circles denote data points obtained from the dephasing of the B(III) and B(IV) units, respectively.

511 and assignment based upon spatial proximity (in the present 512 case between  $^{11}$ B and  $^{31}$ P nuclei), detected by reintroducing 513 (recoupling) magnetic dipole—dipole interactions into MAS 514 NMR experiments via coherent pulse trains. This is done by 515 comparing the signal diminution  $S_0$ —S relative to that ( $S_0$ ) 516 obtained on the sample without recoupling. In the present 517 glasses, heteronuclei occurring within the second coordination 518 sphere from the observed nuclei make the dominant 519 contribution to this diminution. In these experiments, dipolar 520 mixing times of 4.0 ms (60 rotor cycles) have been used. In both 521 samples, the four-coordinated boron species show strong 522 dephasing, while dephasing is significantly weaker for the B(III) 523 units. Based on this finding, we can conclude that only the B(IV) 524 units are involved in B—O—P linkages, whereas B(III) units do 525 not link to phosphate to any significant extent—the slight

dephasing observed here can be easily explained by weaker 526 dipolar interactions with more remote <sup>31</sup>P nuclei. Also, parts a 527 and b of Figure 2 indicate that a significant <sup>11</sup>B{<sup>31</sup>P} REDOR 528 effect is observed for both of the B(IV) signal deconvolution 529 components near 0 and -2.5 ppm, signifying B-O-P linking 530 for both of them. For further quantification, we turn to the 531 REDOR curves (plots of  $\Delta S/S_0$  versus dipolar mixing times  $NT_r$ , 532 see Figure 2c-d). The initial data regime ( $\Delta S/S_0 < 0.2$ ) has been 533 fitted to eq 1 in order to extract dipolar second moments 534  $(M_{2(B-P)})$  for each sample (see Table 3). Here, the very low 535 t3  $M_{2(B-P)}$  value (0.1 ×  $10^{6}$  rad<sup>2</sup>/s<sup>2</sup>) obtained for the B(III) units 536 can be taken as an estimate of a contribution arising from more 537 remote nuclei, which also needs to be accounted for when 538 analyzing the  $M_2$  values measured for the B(IV) units. 539 Accordingly, the corrected  $M_2$  for B(IV) units,  $M_{2(B-P)}$  (B(IV), 540 net), is calculated according to the equation  $M_{2(B-P)}$  (B(IV), 541 net) =  $M_{2(B-P)}$  (B(IV)) –  $M_{2(B-P)}$  (B(III)), and it is evaluated in 542 terms of the number of B(IV)-O-P linkages. Thus, by directly 543 comparing the  $M_{2(B-P)}$  values with that measured for crystalline 544 BPO<sub>4</sub> (four B–O–P linkages at an internuclear distance of 270 545 pm), <sup>28</sup> we can deduce from the information given in Table 3 and 546 Table S4 that the average number of B-O-P linkages per B(IV) 547 species in PB3-P7 is  $\langle m_P(B(IV)) \rangle = 0.37$ . Considering the  $N_4$  548 value, we conclude that, overall, the 60 boron atoms of this glass 549 formulation are involved in 12.5 B-O-P linkages. For the MB3- 550 P5 glass, the average number of B-O-P linkages per B(IV) unit 551 is  $\langle m_p(B(IV)) \rangle = 0.21$ ; in this case, the 50 boron atoms of the 552 glass formulation make a total of 6.7 B-O-P linkages.

**4.3.** <sup>31</sup>P MAS NMR, Refocused INADEQUATE, <sup>31</sup>P{<sup>23</sup>Na} 554 REDOR, and <sup>31</sup>P{<sup>11</sup>B} REAPDOR. Figure 3 shows the <sup>31</sup>P MAS 555 f3 NMR spectra of glasses from series PB2, PB3, MB2, and MB3, 556 while Figure S3 presents the <sup>31</sup>P MAS NMR spectra of glasses in 557 the series PB1, MB1, and PA1-PA3. All the <sup>31</sup>P line shape 558 parameters and the assignment to the various  $P^n_{mX}$  species are 559 summarized in Table S5. The spectra from all the series show 560 three main features: a low intensity peak at ~16 ppm (P<sup>0</sup> units), 561 a strong peak centered between 1 and 4 ppm (P<sup>1</sup> units), and a 562 broader feature between -5 and -10 ppm (P<sup>2</sup> units). The 563 chemical shift range associated with the latter suggests a mix of 564  $P^2_{2B}$  and  $P^2_{1B,1P}$  units, <sup>39</sup> however, connectivity to Si must also be 565 considered. As the concentration of  $P_2O_5$  increases in the PB and 566 MB series, a decrease in the  $P^0$  signal intensity can be observed, 567 while the intensities of the  $P^2$  species increase monotonically. 568

Table 3. Dipolar Second Moments  $(M_{2(S-I)})$  of Glasses and Model Compounds, As Determined from Parabolic Fits of S{I} REDOR Data within  $\Delta S/S_0 \leq 0.20^b$ 

	$M_{2(B-P)} (10^6 \text{ rad}^2/\text{s}^2 (\pm 10\%))$		$\sum r_{\rm (B-P)}^{-6}$	$\sum r_{\rm (B-P)}^{-6} (10^{60} \mathrm{m}^{-6})$		$\langle m_{ m P}({ m B(IV)}) angle \; (\pm 10\%)^a$		
sample ID	B(III)	B(IV)	B(IV), net	B(III)	B(IV)	<sup>11</sup> B{ <sup>31</sup> P} REDOR	<sup>31</sup> P MAS	<sup>31</sup> P{ <sup>11</sup> B} REAPDOR
$BPO_4$	_	_	20.3 (11.2)	_	0.0105	_	_	_
PB3-P7	0.2 (0.1)	2.0 (1.1)	1.8 (1.0)	0.0001	0.0010	0.37	0.42	0.41
MB3-P5	0.2 (0.1)	1.3 (0.7)	1.1 (0.6)	0.0001	0.0007	0.21	0.26	0.31
sample ID	$M_{2({\rm Na-P})} (10^6 {\rm rad}^2/{\rm s}^2)$		$\sum r_{(1)}$	$_{\text{Na-P})}^{-6} (10^{60} \text{ m}^{-6})$		$M_{2(P-Na)} (10^6 \text{ rad}^2/\text{s}^2)$	Σ	$r_{\rm (P-Na)}^{-6} (10^{60} \text{ m}^{-6})$
$Na_3P_3O_9$	5.4 (3.4) - 0.3 (0.2) 1.1 (0.7) 2.3 (1.4)			0.0041 —		26.7 (8.2)	26.7 (8.2) 0.0041	
PB0						24.4 (7.5) 0.003		_
PB3-P1			0.0002		0.0042			
PB3-P4			0.0008		0.0037			
PB3-P7			0.0018		0.0035			

"Average number of B–O–P linkages per B(IV) unit. <sup>b</sup>Here, S and I denote the observed and the non-observed nuclei.  $M_{2(S-I)}$  values in parentheses are raw data before model compound calibration.  $M_{2(B-P)}$  data listed for B(IV), net and B(IV), respectively, refer to values with and without correction from the contribution of more remote <sup>11</sup>B nuclei to the REDOR curvature. Additionally, calculated values of the average number of P next nearest neighbors around B(IV) were calculated from both double resonance techniques and <sup>31</sup>P MAS NMR speciation.

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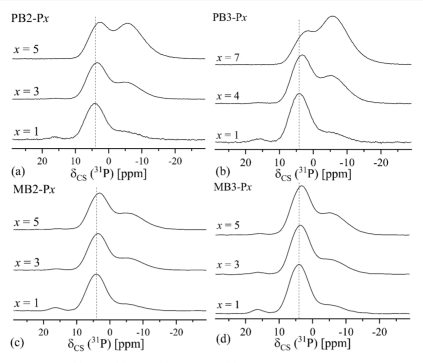


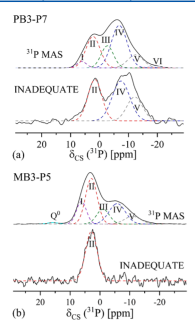
Figure 3. <sup>31</sup>P MAS NMR spectra of series (a) PB2, (b) PB3, (c) MB2, and (d) MB3 glasses.

Peralkaline glasses show a slight increase in the intensity of 569 that peak as the concentration of P<sub>2</sub>O<sub>5</sub> increases from 1 to 3 mol %, while the changes in the P<sup>0</sup> signal intensity depend upon the substitution scheme used (see Figure S3). The spectra of glasses belonging to the series PA1 show a narrowing of the P<sup>0</sup> and P<sup>1</sup> resonance lines while going from PA1-P1 to PA1-P3 in addition to a low-frequency displacement and increased intensity of the peak. These changes may indicate the development of more 577 highly ordered phosphate environments. XRD patterns of the x 5 glasses show the formation of crystalline sodium pyrophosphate (Na<sub>4</sub>P<sub>2</sub>O<sub>7</sub>), a small amount of which appears 580 to also be present in PA1-P3 glass (see Figure S1 for XRD 581 patterns of PA1-P3 and PA1-P5 samples). Series PA2 and PA3, 582 on the other hand, show similar or slightly decreased P<sup>0</sup> peak intensities from x = 1 to 3, alongside small low-frequency shifts 584 in peak position.

Additionally, in glasses of all regimes and series, the P1 585 586 resonance center shifts toward lower frequencies with increasing 587 P content (as displayed in Figure 3). To aid in the assignments of these signals to the various  $P_{mX}^n$  units, we have also computed the NMR parameters of the phosphate species found in three 590 replicas of models containing about 400 atoms of the PB2-P5 glass (whose data is reported in Table 2). Additional chemical 592 shift calculations have been performed on phosphate and borophosphate compositions (mol %): 55Na<sub>2</sub>O-45P<sub>2</sub>O<sub>5</sub> and  $40\text{Na}_2\text{O} - 18\text{ B}_2\text{O}_3 - 42\text{P}_2\text{O}_5$ . Our results show that  $P^1_{1B(IV)}$ ,  $P^1_{1Si}$ and P<sub>1P</sub> species resonate at about 8.2, 4.3, and 3.8 ppm with standard deviations between 2 and 5 ppm, indicating that these units may not be clearly distinguishable in the broadened MAS NMR spectra. The predicted chemical shifts for the four possible  $P^2$  species, namely  $P^2_{2B(IV)}$ ,  $P^2_{1B(IV),1Si}$ ,  $P^2_{2Si}$  and  $P^2_{1B(IV),1P}$ , are -3.5, -11.5, -10.8, and -14.8 ppm, respectively, while no value 601 for P<sup>2</sup><sub>1Si,1P</sub> can be found as it has not been generated in the MD 602 simulation of the phospho-borosilicate glass. To further 603 understand the compositional evolution of these spectra, various 604 advanced NMR experiments have been performed to recouple 605 homo- and heteronuclear spin-spin interactions, thereby providing direct evidence of P-O-P and P-O-B connectiv- 606 ities.

1-D refocused INADEQUATE pulse sequences have been 608 applied to probe the indirect <sup>31</sup>P-<sup>31</sup>P spin-spin interactions in 609 the samples PB3-P7 and MB3-P5. As this method relies on the 610 existence of through-bond spin-spin interactions to generate 611 double quantum coherences, it can serve as a filter for selectively 612 detecting only those <sup>31</sup>P nuclei that are involved in P-O-P <sub>613</sub> linkages. Part a and b of Figure 4 show direct comparisons 614 f4 between the <sup>31</sup>P MAS NMR and the INADEQUATE spectra for 615 two selected glass compositions. The double-quantum filtered 616 spectrum of MB3-P5 shows a single component near 2.5 ppm. 617 The same peak is also identified in the INADEQUATE 618 spectrum of the sample PB3-P7. However, an additional 619 component is observed near -10 ppm, whereas the difference 620 spectrum shows components near +4.6 and -4.6 ppm. Using the 621 line shape parameters of these partial spectra as additional 622 constraints, the overall <sup>31</sup>P MAS NMR spectrum can be fitted to 623 a total of five Gaussian contributions I-V, whose parameters are 624 summarized in Table S5 (see Figure 4). This particular 625 deconvolution model has been used to fit the <sup>31</sup>P MAS NMR 626 spectra of all the glasses by maintaining similar Gaussian line 627 widths and positions, allowing the fwhm and  $\delta_{\mathrm{CS}}^{\mathrm{iso}}$  to vary within 628 the constraints ±0.1 ppm and ±0.5 ppm, respectively. The lone 629 exception to the fitting methodology concerned the PA1-P3 630 glass composition which, as mentioned previously, exhibited 631 some degree of crystallinity marked by a significant reduction in 632 the fwhm of component II, whose chemical shift agrees with that 633 of pyrophosphate  $(P_{1P}^1)$  units. We further note that component 634 II in all samples is narrower than the overall signal near 1-4 ppm 635 observed in the single-pulse spectra, requiring the additional 636 components I and III for a satisfactory deconvolution. On the 637 basis of the <sup>31</sup>P{<sup>11</sup>B} REAPDOR results detailed below, 638 component I is tentatively assigned to P11B units. For 639 component III, the possibilities include  $P_{1Si}^1$  and  $P_{2B}^2$ .

Further insight into this question comes from  $^{31}P\{^{11}B\}$  641 REAPDOR experiments. Parts a and b of Figure 5 show the 642 f5



**Figure 4.** Deconvolution of the <sup>31</sup>P MAS NMR and double-quantum filtered <sup>31</sup>P MAS NMR spectra, using the refocused INADEQUATE sequence: (a) PB3-P7 and (b) MB3-P5 samples. The double-quantum filtered spectra were utilized to guide the fitting of <sup>31</sup>P MAS NMR spectra. Red, blue, and green data points represent the full spectra, the filtered spectra, and the difference spectra, respectively.

643 Fourier transforms of rotor-synchronized  $^{31}P$  spin echoes 644 without  $(S_0)$  and with (S)  $^{11}B$  dipolar recoupling for the 645 samples PB3-P7 and MB3-P5 after 16 and 22 rotor cycles. Also

displayed are the difference spectra  $(S_0 - S)$ , which indicate that 646 the phosphate species contributing to both major resonances 647 near 1-4 ppm and -5 to -10 ppm show significant  ${}^{11}B-{}^{31}P$  648 dipole-dipole coupling. We conclude that these phosphate 649 species have, at least in part, <sup>11</sup>B next nearest neighbors. Two and 650 three points on the REAPDOR curve have been measured for 651 MB3-P5 and PB3-P7, respectively, and they are compared with 652 data generated from SIMPSON simulations taking into account 653 the natural abundance of the <sup>11</sup>B isotope (80.4%) (see Figure 654  $5c_{r}d$ ). In these simulations, the P-B distance across a P-O-B 655 linkage has been assumed to be 270 pm, consistent with the 656 shortest distance in BPO<sub>4</sub>. 64 Simulations are based on a 11B 657 quadrupolar coupling constant of 0.52 and 0.64 MHz for glasses 658 MB3-P5 and PB3-P7, respectively. These values have been taken 659 from the overall spectral range of the spinning sideband pattern 660 associated with the  $|m| = \frac{1}{2} \leftrightarrow |m| = \frac{3}{2}$  satellite transitions (see 661 Figure S4). For spin-3/2 nuclei, this overall spectral range is 662 identical with the quadrupolar coupling constant,  $C_{\rm O}$ . We can 663 see from the simulations that the signal near -5 to -10 ppm 664 shows a stronger REAPDOR effect compared to that simulated 665 for a single P-O-B linkage (about 1.1 and 1.2 per P unit in PB3- 666 P7 and MB3-P5), whereas the signal in the 1 to 4 ppm range 667 shows a weaker interaction (about 0.65 and 0.8 P-O-B 668 linkages per P unit). While the P<sup>0</sup> peak also shows evidence of 669 some weak dephasing in the MB3-P5 sample, this effect must be 670 attributed to contributions from more remote <sup>11</sup>B nuclei in the <sup>671</sup>

In light of the above-discussed results, the most likely 673 candidates for structural species detected in our experiments, 674 based upon the studied compositional regime and the 675 experimental  $^{31}P$  chemical shifts, are  $P^{1}_{1B(IV)}$  (component I), 676

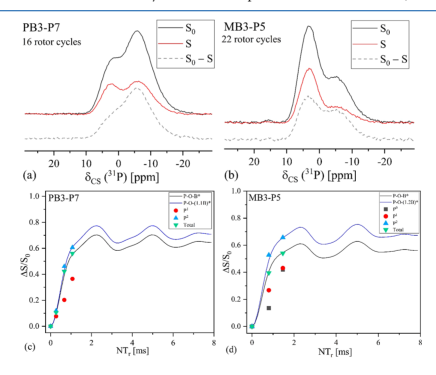


Figure 5. Fourier transforms of the <sup>31</sup>P MAS spin echoes obtained in a <sup>31</sup>P{<sup>11</sup>B} REAPDOR experiment: reference signal,  $S_0$ , signal with dipolar dephasing, and REAPDOR difference spectra  $S_0$  – S. Key: (a) PB3-P7 glass after 16 rotor cycles ( $NT_r$  = 1.1 ms) and (b) MB3-P5 glass after 22 rotor cycles ( $NT_r$  = 1.5 ms). (c and d) <sup>31</sup>P{<sup>11</sup>B} REAPDOR curves ( $\Delta S/S_0$  vs  $NT_r$ ) for PB3-P7 and MB3-P5 samples, respectively, for the resolved resonances of P<sup>0</sup>, P<sup>1</sup>, and P<sup>2</sup> units and the integral, after a given evolution time. The solid lines represent the results from numerical simulations made with the SIMPSON software. The black curve represents the REAPDOR curves for a <sup>11</sup>B–<sup>31</sup>P two-spin system assuming an internuclear distance equal to that of a P–O–B linkage found in BPO<sub>4</sub> (270 pm). The blue curve has been scaled to the experimental data of the P<sup>2</sup> species in order to deduce an average number of P–O–B linkages.

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Table 4. Average Number of B Next Nearest Neighbors around P and Average Number of Bridging Oxygen Atoms per Phosphate Unit As Determined from Fitted <sup>31</sup>P MAS NMR Data<sup>c</sup>

	<sup>31</sup> P MAS NMR		<sup>29</sup> Si MA	S NMR	<sup>23</sup> Na MA	<sup>23</sup> Na MAS NMR	
sample ID	$\langle m_{\rm B}({ m P}) \rangle$	$\langle n \rangle$	Si <sup>4</sup> (calcd)	$\delta_{ m cg}$	$\delta_{ ext{CS}}^{ ext{iso}}  ext{(ppm)}$	C <sub>Q</sub> (MHz	
PB0	-	-	70.8	-91.9	-5.2	2.4	
PB1-P1	0.50	1.28	83.4	n.d. <sup>b</sup>	-5.2	2.5	
PB1-P3	0.73	1.56	92.0	n.d.	-5.8	2.5	
PB1-P5	0.91	1.77	97.7	n.d.	-5.4	2.6	
PB2-P1	0.47	1.23	81.5	n.d.	-5.3	2.4	
PB2-P3	0.65	1.45	84.5	n.d.	-5.0	2.6	
PB2-P5	0.90	1.74	88.3	n.d.	-5.7	2.5	
PB3-P1	0.48	1.24	80.8	-92.3	-5.9	2.2	
PB3-P4	0.73	1.50	95.1	-95.3	-6.5	2.2	
PB3-P7	$1.02/1.20^a$	1.83/1.86 <sup>a</sup>	100.3	-99.7	-6.4	2.2	
MB0	_	_	66.7	-92.1	-3.0	2.7	
MB1-P1	0.72	1.29	77.4	-93.0	-6.4	2.9	
MB1-P3	0.84	1.50	88.1	-96.2	-7.9	2.8	
MB1-P4	0.88	1.56	91.4	-97.5	-8.8	2.8	
MB2-P1	0.69	1.25	74.6	-92.5	-6.0	2.9	
MB2-P3	0.81	1.45	77.8	-95.2	-7.8	2.8	
MB2-P5	0.86	1.56	92.1	-97.4	-9.0	2.7	
MB3-P1	0.69	1.24	74.1	-91.9	-5.8	2.8	
MB3-P3	0.80	1.42	85.1	-94.1	-7.3	2.8	
MB3-P5	0.84	1.55	92.3	-96.7	-9.0	2.7	
PA0	_	_	61.7	-91.3	-4.8	2.9	
PA1-P1	0.61	1.21	68.9	-94.6	-6.7	2.8	
PA1-P3	0.62	1.30	84.0	-96.4	-6.8	2.8	
PA2-P1	0.67	1.24	68.3	-92.8	-5.1	2.9	
PA2-P3	0.74	1.39	77.3	-95.2	-7.0	2.8	
PA3-P1	0.66	1.21	66.6	-92.7	-5.0	2.9	
PA3-P3	0.71	1.33	77.8	-95.1	-6.2	2.8	

<sup>a</sup>Two separate sets of measurements with fits constrained by refocused INADEQUATE experiments. <sup>b</sup>Not determined. <sup>c29</sup>Si chemical shifts  $\delta_{cg}$  (center of gravity,  $\pm$  0.5 ppm), calculated Si<sup>4</sup> fractions ( $\pm$ 1.0%) based on charge balance, average <sup>23</sup>Na isotropic chemical shifts,  $\delta_{CS}$  ( $\pm$  0.5 ppm), and quadrupolar coupling constants  $C_Q$  ( $\pm$  0.2 MHz).

677  $P_{1P}^1$  (component II), and  $P_{2B}^2$  (component III). Additionally, 678 P<sup>1</sup><sub>1Si</sub> species may contribute to component II or III, which in the 679 latter case would have the effect of weakening the overall 680 REAPDOR response. Plausible assignments for component IV and V are the species  $P^2_{1B,1P}$  and  $P^3_{2B,1P}$  as these signals appear 682 both in the refocused INADEQUATE experiments and in the 683 REAPDOR difference spectra. 38 In addition, subtle differences 684 arising from a detailed comparison of the refocused INAD-685 EQUATE spectra and the REAPDOR difference spectra suggest that the P<sup>1</sup><sub>1Si</sub> and P<sup>2</sup><sub>1B.1Si</sub> units may be present as well, the former contributing intensity mostly to signals III and IV and latter contributing intensity mostly to signal V in the sample PB3-P7. Concerning the sample MB3-P5, we further believe that the resonances associated with P<sup>2</sup><sub>1B,1P</sub> and P<sup>3</sup><sub>2B,1P</sub> units may have eluded the double-quantum filtered detection in Figure 4b due to their low concentrations. Minor components labeled P<sup>0</sup> and VI are attributed to isolated monophosphate  $(P^0)$  or  $P^2_{2P}$  chain phosphate species which do not contain any linkages to boron. Table S5 presents the proposed phosphate speciation of all the samples as determined from the spectral deconvolutions. In the case of sample PB3-P7, two separate sets of analyses including refocused INADEQUATE experiments have been performed; the deviations give an impression of the possible experimental error in the speciation.

The data presented in Table S5 reveals some universal trends, roz regardless of R regime: As x increases from 1 to higher values, the roz concentrations of  $P^0$  and  $P^1$  species decrease monotonically,

while those of  $P^2$  and  $P^3$  increase. The concomitant increase in 704 connectivity can be expressed by the average number of bridging 705 oxygen atoms per phosphate unit given by eq 3 and presented in 706 Table 4.

$$\langle n \rangle = 0 \times f(P^0) + 1 \times f(P^1) + 2 \times f(P^2) + 3 \times f(P^3)$$
(3) 708

Furthermore, for all the glass compositions investigated in the 709 present study, a general rise in P-O-B linked phosphate units 710 has been observed with increased P2O5 presence in the glass 711 network, with a preference toward units that have at least two 712 bridging oxygen atoms (P<sup>2</sup> units). While there is a decrease in 713 the concentration of  $P_{1B}^1$  units, those of other B linked  $P_{1B}^2$  units 714  $(P_{2B}^2, P_{1B,1Si}^2, P_{1B,1P}^2)$  and some  $P_{2B,1P}^3$  units as well become 715 more prominent, with the average number of B next nearest 716 neighbors for each P in the glass rising from 0.5 to 0.9-1.2 in PB 717 glasses, rising from 0.7 to 0.8-0.9 in MB glasses, and either 718 remaining similar or rising from 0.6 to 0.7 in PA glasses (Table 719 4). These values have been calculated from the  $P^{n}_{mB}$  720 concentrations, according to the assessed speciation from <sup>31</sup>P <sub>721</sub> MAS NMR. While in glasses with x = 1 about half (40–60%) of 722 the phosphate units have no linkages to boron (i.e., are either P<sup>0</sup> 723 or P<sub>1P</sub>, higher concentrations of P<sub>2</sub>O<sub>5</sub> are linked more 724 extensively with B, as they contain higher levels of P2 units. 725 This effect is most pronounced in the perboric (PB) glass having 726 the highest  $P_2O_5$  content (e.g., PB3-P7).

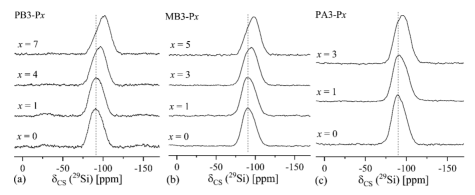


Figure 6. <sup>29</sup>Si MAS NMR spectra of series (a) PB3, (b) MB3, and (c) PA3 glasses. The dashed lines are guides to the eye.

To validate our results, we can compare the average dipolar results interaction strengths measured in the  $^{11}\mathrm{B}\{^{31}\mathrm{P}\}$  REDOR and the results are consistent results, i.e., the total number of B–O–P linkages as determined from results are determined from results as determined from results as determined from results are results. The total number of B–O–B linkages derived from results are results are results. The results are results are results are results are results. The results are results are results are results are results are results are results. The results are results are results are results are results are results are results. The results are results. The results are results ar

$$\#(B - O - P) = N_4 \times 2 \times \text{mol. } \%B_2O_3 \times \langle m_P(B) \rangle$$
 (4)

739 As detailed above, 60 boron atoms in glass PB3-P7 make a total 740 of 12.5 B-O-P linkages whereas the 50 boron atoms in glass 741 MB3-P5 make a total of 6.7 B-O-P linkages.

738

For comparison, the average number of B-O-P linkages per can be deduced from the simulation of the REAPDOR 744 dephasing observed for the total integrated <sup>31</sup>P MAS NMR 745 signal (see Figure 5, parts a and b). As shown in Figure 5c-d, this 746 number is found near  $\langle n_{\rm B}({\rm P}) \rangle = 1.0$  for both samples. Thus, the 747 total number of P-O-B linkages is given by #(P-O-B) =748  $2x\langle n_B(P)\rangle$ . For example, the 14 phosphorus atoms in glass PB3-749 P7 make a total of 14 P-O-B linkages whereas the 10 750 phosphorus atoms in glass MB3-P5 make a total of 10 P-O-B 751 linkages. Yet another independent estimate of #(P-O-B) is 752 available from the phosphorus speciation as listed in Table S5. 753 From the calculated values  $\langle n_B(P) \rangle$  shown in Table 4, we 754 conclude that there are 14.3/16.8 and 8.6 P-O-B linkages in 755 samples PB3-P7 (measured twice) and MB3-P5, respectively, in 756 good agreement with the REAPDOR estimate. This consistency 757 also confirms our <sup>31</sup>P MAS NMR peak assignments within the entire body of NMR data collected. Table 3 displays these 759 comparisons for the two studied samples, expressing the results 760 in terms of the average number of B-O-P linkages per B(IV) 761 unit, as calculated from the phosphorus speciation according to 762 eq 5.

$$\frac{\text{avg. } \#\text{B-O-P linkages}}{\text{B(IV)}}$$

$$= \frac{(\text{avg. } \#\text{B per P}) \times (\text{P content, mol. }\%)}{(N_4 \text{ fraction}) \times (\text{B content, mol. }\%)}$$
(5)

764 All the results have been summarized in Table 4. The data 765 presented in Table 4 further shows that the rate of change of 766  $\langle n_{\rm B}({\rm P}) \rangle$  as a function of x increases in the order PA  $\rightarrow$  MB  $\rightarrow$  767 PB, as does the number of P-O-B linkages formed for a fixed x-768 value. Thus, the more highly polymerized PB glasses tend to mix 769 more extensively with P than the MB or PA glasses as the P<sub>2</sub>O<sub>5</sub>

content is increased. However, due to the well-known affinity of 770 phosphate units toward sodium, the extent of P-O-B mixing is 771 also highly dependent upon the Na $^+$  distribution in the network. 772 Alternatively, the affinity of glasses to form P-O-B(IV) 773 linkages can also greatly affect Na $^+$  distribution among NFUs, 774 as the average bond valence of tetrahedral phosphate and boron 775 units balance  $\left(\frac{1.25+0.75}{2}=1\right)$ , 86 indicative of high bond stability 776 and presumptive charge-sharing between negatively charged 777 B(IV) units and nonbridging oxygen on phosphate species. 87,88 778 These competing factors will be further discussed in the next 779 sections.

**4.4.** <sup>29</sup>Si MAS NMR. Figures 6 and S5 present the <sup>29</sup>Si MAS 781 f6 NMR spectra of samples in the series MB3, PB3, and PA3 as well 782 as MB1, MB2, PA1, and PA2 samples. Each series displays a 783 broad peak centered near -92 ppm in the phosphate-free glass 784 which shifts significantly toward lower frequency with increased 785 P<sub>2</sub>O<sub>5</sub> content. A similar trend has also been observed by Muñoz 786 et al.<sup>39</sup> in a compositionally related borosilicate system. The 787 signal positions we see in our samples are also consistent with 788 previously studied P2O5-free and P2O5-containing alkali 789 borosilicate glasses of similar R and K (=  $[SiO_2]/[B_2O_3]$ ) 790 values. 39,89,90 Owing to the poor resolution, no unambiguous 791 line shape deconvolution is possible here. Therefore, Table 4 792 lists the average <sup>29</sup>Si chemical shifts determined from the center 793 of gravity. We clearly note that this quantity shifts significantly 794 toward lower frequency with increasing P2O5 content. As the 795 <sup>29</sup>Si chemical shifts in compositionally complex mixed-network 796 former glasses are influenced by numerous different factors, 797 including the number of NBOs and the number of next nearest 798 neighbor linkages the trend observed here can be explained in 799 different ways. Further guidance comes from quantum chemical 800 calculations of chemical shifts done on PB2-P5 glass, which are 801 summarized in Table 2 for various Si<sup>4</sup> species with different 802 connectivities. The calculations confirm the trends found by 803 Fortino et al.:<sup>71</sup> While  $\delta_{\rm CS}^{\rm iso}(^{29}{
m Si})$  increases when Si atoms are  $_{804}$ replaced by borate species in the second coordination sphere, 805 the effect is significantly stronger for Si-O-B(IV) linkages than 806 for Si-O-B(III) linkages. For example, the computed isotropic 807 chemical shift for a  $Si_{4B(IV)}^4$  species is -82.0 ppm while it is 808-107.6 ppm for a  $Si_{4B(III)}^4$  species<sup>71</sup> (see Table 2). Intermediate 809 values are found for Si<sup>4</sup> species with mixed B(III), B(IV), and Si<sup>4</sup> 810 ligation. The chemical shift ranges of such species overlap with 811 those usually associated with Si<sup>2</sup> and Si<sup>3</sup> units in silicate glasses, 812 making a corresponding assignment difficult. The situation is 813 complicated further if ligation to phosphorus is considered, as 814 replacement of a Si-O-Si linkage by a Si-O-P linkage 815 produces a shift that is more negative than that of Si<sup>4</sup><sub>4Si</sub>. Given 816

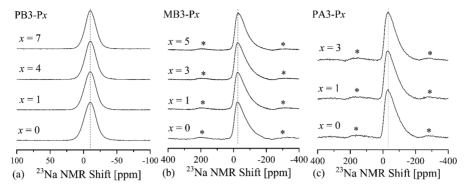
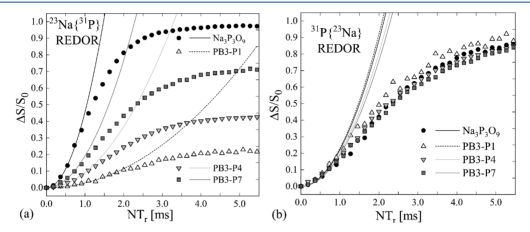


Figure 7. <sup>23</sup>Na MAS NMR spectra of series (a) PB3, (b) MB3, and (c) PA3 glasses. Series PB3 spectra were collected at 16.4 T while series MB3 and PA3 spectra were collected at 5.7 T. The asterisks mark spinning sidebands and the dashed lines serve as guides to the eye.



**Figure 8.** (a)  $^{23}$ Na $\{^{31}$ P $\}$  and (b)  $^{31}$ P $\{^{23}$ Na $\}$  REDOR curves for the model compound Na $_3$ P $_3$ O $_9$  and the glass samples PB0, PB3-P1, PB3-P4, and PB3-P7. The solid curves represent parabolic fits to the initial regime ( $\Delta S/S_0 \le 0.20$ ) from which the heteronuclear second moments are extracted via eq 1.

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817 this situation and the poorly resolved lineshapes, it is not 818 possible to deduce detailed silicon speciation from the  $^{29}$ Si MAS 819 NMR spectra alone. Si<sup>4</sup> species connected to a mix of Si, B(III), 820 and B(IV) are likely to be present; in addition, there may be 821 some contribution from Si<sup>3</sup> units. In similar sodium phospho-822 borosilicate glasses, Muñoz et al. Si attributed the Si chemical 823 shift trend observed upon the addition of  $P_2O_5$  to a decrease in 824 silicate-borate network connectivity. However, it might also 825 reflect a diminution of Si units or, in principle, an increase in the 826 number of Si-O-P linkages. From the charge balance 827 constraint (eq 6),

$$[Si^3] + [B(IV)] + [P^2] + 2[P^1] = [Na]$$
 (6)

829 we can deduce the concentration of  $\mathrm{Si}^3$  units if we neglect the 830  $\mathrm{B}^2(\mathrm{III})$  units. These numbers, listed in Table 4, show that the 831 addition of phosphate in these glasses tends to diminish [Si³], 832 i.e., the concentration of NBOs in the silicate network. Again this 833 effect can be attributed to the successful competition of  $\mathrm{P}_2\mathrm{O}_5$  834 over  $\mathrm{SiO}_2$  in attracting network modifier and/or effective 835 charge-sharing between P and B units in P-O-B(IV) linkages 836 in the absence of network modifying species.

4.5. <sup>23</sup>Na MAS NMR and <sup>23</sup>Na{<sup>31</sup>P} and <sup>31</sup>P{<sup>23</sup>Na} REDOR. Figure 7a—c presents the <sup>23</sup>Na MAS NMR spectra of series MB3, PB3, and PA3. The differences in appearance are attributed to the different magnetic field strengths used for data collection. The spectra for MB and PA samples have been collected at 5.7 T, while PB samples have been measured at 16.4 T. Therefore, the extent of second-order quadrupolar broades ening is very different for the low and high field data. The <sup>23</sup>Na

isotropic chemical shift ( $\delta_{
m CS}^{
m iso}$ ) and  $C_{
m Q}$  values listed in Table 4 845 have been extracted from the fitted spectra using the Czjzek 846 distribution model implemented in DMfit. The isotropic 847 chemical shift values in MB glasses show a monotonic decrease 848 from near -3 to -9 ppm with increasing x, while PB and PA 849 glasses show much less pronounced effects. The isotropic 850 chemical shift is expected to be highly dependent upon the 851 distribution of sodium among the silicate, borate, and phosphate 852 structural units, whose anionic charges it compensates. On the 853 basis of the findings of the present study, the main change as a 854 function of x in all three series is the increased participation of 855 phosphate in the anionic inventory of the glass. The overall <sup>23</sup>Na 856 chemical shift trend toward more negative values with increasing 857 x reflects the increased importance of sodium/phosphate 858 interactions, and the simultaneous decrease of the importance 859 of B(IV)-Na and Si<sup>3</sup>-Na interactions.

Further, to probe Na–P interactions in the glass network, 861  $^{23}$ Na $\{^{31}$ P $\}$ , and  $^{31}$ P $\{^{23}$ Na $\}$  dipole—dipole couplings have been 862 measured in the glasses and the crystalline model compound 863 Na<sub>3</sub>P<sub>3</sub>O<sub>9</sub> for calibration. Again,  $M_2$  values have been extracted 864 via eq 1 from the initial portion of each REDOR curve ( $\Delta S/S_0 < 865$  0.2; see Figure 8), and are presented in Table 3. As anticipated, 866 fs  $M_{2(Na-P)}$  increases with increasing x. We estimate the average 867 coordination of the sodium ions with phosphorus in the second 868 coordination sphere at 0.3, 1.0, and 2.1 P for samples PB3-P1, 869 PB3-P4, and PB3-P7, respectively. This trend in  $^{23}$ Na $^{-31}$ P  $^{870}$  proximity is accompanied by decreases in the Si $^3$  and  $N_4$   $^{871}$  fractions (see Tables 4 and S4), suggesting that Na $^+$  ions,  $^{872}$  which in the phosphate-free borosilicate glasses either charge

Table 5.  $N_4$  Values and Fractions of B<sup>4</sup>(IV), B<sup>3</sup>(IV), B<sup>3</sup>(III), and B<sup>2</sup>(III) Units Found in the MD-Generated Structural Models of the Investigated Glasses and Average Numbers of Network Former Cations Bound to Fully Connected B<sup>4</sup>(IV) and B<sup>3</sup>(III) Units<sup>a</sup>

sample ID	$N_4$	$B^3(IV)$	$B^4(IV)$	#B(IV)	#B(III)	#Si	#P	#P ( <sup>31</sup> P MA	S NMR)
PB0	59.5 [62.9]	2.0	57.5	1.0	1.0	2.0	_	_	
PB2-P1	60.5 [63.5]	1.6	58.9	0.9	1.0	2.1	0.0	0.0	)
PB2-P3	58.6 [62.1]	1.8	56.8	0.9	1.1	1.9	0.2	0.3	l
PB2-P5	54.6 [59.1]	1.3	53.3	0.7	1.1	1.9	0.3	0.3	3
PB3-P7	53.8 [56.4]	3.9	49.9	0.7	1.1	1.7	0.5	0.4	1
MB0	61.7 [67.2]	4.4	57.3	0.9	0.8	2.3	_	_	
MB2-P1	63.4 [67.5]	3.4	60.0	1.0	0.9	2.1	0.0	0.0	)
MB2-P3	60.9 [67.2]	2.7	58.2	0.8	0.9	2.2	0.2	0.2	2
MB2-P5	60.3 [65.5]	2.2	58.1	0.7	0.9	2.2	0.3	0.3	3
PA0	67.2 [71.0]	5.8	61.4	0.8	0.6	2.7	-	-	
PA2-P1	58.9 [71.4]	2.1	56.8	0.6	0.8	2.5	0.0	0.0	)
PA2-P3	66.6 [71.9]	3.4	63.2	0.8	0.6	2.5	0.2	0.2	2
Sample ID	$N_3$	$B^2(III)$	В	$3^3(III)$	#B(IV)	#B(III)		#Si	#P
PB0	40.5	9.9		30.6	1.7	0.3		1.0	_
PB2-P1	39.5	6.7		32.8	1.6	0.4		1.0	0.0
PB2-P3	41.4	4.1		37.3	1.7	0.3		1.1	0.0
PB2-P5	45.4	4.5		40.9	1.5	0.4		1.1	0.0
PB3-P7	46.2	1.7		44.5	1.5	0.5		1.0	0.0
MB0	38.3	12.5		25.8	1.6	0.3		1.1	_
MB2-P1	36.6	7.5		29.1	1.6	0.2		1.2	0.0
MB2-P3	39.1	7.4		31.7	1.5	0.3		1.2	0.0
MB2-P5	39.7	4.7		35.0	1.4	0.3		1.2	0.1
PA0	32.8	13.9		18.9	1.5	0.1		1.4	-
PA2-P1	41.1	10.0		31.1	1.4	0.2		1.4	0.0
PA2-P3	33.4	8.6		24.8	1.4	0.2		1.4	0.0

<sup>&</sup>quot;# indicates the average number of linkages from  $B^4(IV)$  and  $B^3(III)$  species to each NFU.  $N_4$  values measured by NMR are listed in brackets. #P neighbors around  $B^4(IV)$  have also been compared to the values as calculated from fitting of  $^{31}P$  MAS NMR data.

874 compensate B(IV) units or create NBOs in the silicate network, 875 are now more closely associated with phosphate units. 876 Consistent with this interpretation, the  $M_{2(P-Na)}$  values 877 measured in the reverse  $^{31}P\{^{23}Na\}$  REDOR experiments remain 878 roughly constant as a function of  $\alpha$ , suggesting a rather constant 879 environment of the phosphate units with sodium ions.

4.6. MD Simulations: Glass Structure Predictions. MD simulations have been performed on one series per Na/B regime series PB2, MB2, and PA2; including the  $P_2O_5$ -free countersparts) in order to (i) compare and prove the validity of using MD simulations to reproduce and predict glass structures and sets (ii) obtain additional structural details that are not readily see available from MAS NMR.

Boron Environment: Short-Range Order. Reliable B-O interatomic parameters must first be validated by their ability to 889 accurately reproduce the  $N_4$  values as a function of composition. In this respect, Table 5 indicates good agreement between simulated and experimental data. For each series investigated, 892 the majority of the boron atoms are four-coordinated ( $N_4$  > .53), although the experimental  $N_4$  values are always higher by -7% compared to the simulated data. Nevertheless, the simulations reflect the same trends observed experimentally:  $N_4$ tends to decrease in the order  $PA \rightarrow MB \rightarrow PB$  and with increasing P<sub>2</sub>O<sub>5</sub> content. Table 5 further indicates that the large majority (>91%) of the four-coordinated boron species B(IV) are bound to four bridging oxygen atoms, i.e., they are B<sup>4</sup>(IV) 900 species. The fraction of B(IV) species linked to NBOs, denoted 901 B<sup>3</sup>(IV) species, is negligible. It should be noted here that this 902 kind of information is not available from solid-state NMR, as the 903 NMR-parameters of four-coordinated boron atoms bound to

NBOs are unknown and the spectra of both species are most 904 likely overlapping.

The same kind of analysis has also been conducted on the 906 three-coordinated boron (B(III)) species. Table 5 reveals that 907 the majority of them are B<sup>3</sup>(III) species (trigonal B unit with 908 three BO), but the simulations also suggest that there are non- 909 negligible amounts of B<sup>2</sup>(III) units bearing one nonbridging 910 oxygen. In this context, it is important to note that small 911 concentrations of B2(III) units are difficult to detect and 912 quantify by high-field <sup>11</sup>B MAS NMR, and the low-field data by 913 Yun and Bray based on their detailed analysis of <sup>11</sup>B and <sup>10</sup>B <sub>914</sub> wide-line NMR spectra<sup>33</sup> remain the best estimates known to 915 date. In line with these wide-line NMR results, the MD 916 simulations indicate that the fraction of B<sup>2</sup>(III) units decreases 917 with decreasing R values, i.e., in the order PA  $\rightarrow$  MB  $\rightarrow$  PB. 918 Furthermore, in agreement with these general trends, Table 5 919 reveals that the successive incorporation of P<sub>2</sub>O<sub>5</sub> into these 920 glasses has the effect of depleting the B<sup>2</sup>(III) units drastically. 921 Such information is currently not available experimentally from 922 the <sup>11</sup>B NMR data presented here, even when analyzing boron 923 speciation with <sup>11</sup>B 3QMAS NMR spectroscopy (data not 924

Boron Environment: Connectivity. Table 5 lists the average 926 number of each NFU around each  $B^4(IV)$  and  $B^3(III)$  unit, as 927 extracted from the MD generated models. In general,  $B^4(IV)$  928 species are connected to 1.9–2.3 Si atoms and nearly equal 929 amounts (between 0.7 and 1.1) of B(III) and B(IV) species in 930 the PB2 and MB2 series. In both series, the number of P atoms 931 connected to  $B^4(IV)$  species increases (up to 0.3 per  $B^4(IV)$  932 species) with the addition of  $P_2O_5$ . While the experimental data 933 from NMR do not permit such an incisive numerical analysis, we 934

935 can compare the simulated average number of B(IV)-O-P 936 linkages per boron with the results from  $^{11}B\{^{31}P\}$  REDOR. For 937 the glass PB3-P7, we note an excellent quantitative agreement. 938 Also, the MD simulations agree with the experimental REDOR 939 result that there are very few if any B(III)-O-P linkages. 940 Interestingly, we also see that P prefers to replace B(IV) and Si as 941 opposed to B(III) species in the second coordination sphere of 942  $B^4(IV)$ . In the PA2 series, the second coordination sphere of 943  $B^4(IV)$  has more Si atoms (2.5-2.7) and fewer B(IV) and B(III) 944 units (0.6-0.8) compared to the PB2 and MB2 series. In this 945 case, P seems to preferentially replace Si instead of B(IV). Again, 946 such detailed information is not currently available from solid-947 state NMR.

Overall, the MD simulations confirm the NMR result that 949 phosphorus promotes a restructuring of the borate component. 950  $B^4(IV)$  species are mainly surrounded by silicon atoms and 951 phosphate units, whereas  $B^3(III)$  species are mainly connected 952 to B(IV) species (between 1.4 and 1.7), silicon (1.0–1.4), and to 953 a very small extent, other B(III) species (0.1–0.4). Unlike B(IV) 954 species, B(III) species tend to avoid linking to phosphorus. 955 While the MD simulations suggest that the fraction of B(III) 956 O—Si linkages tends to increase with  $\alpha$ , no such information can 957 be drawn from the NMR data, as they do not reveal any 958 discernible spectroscopic trends for the B(III) units.

The effect of the network former mixing in the second coordination sphere of the boron atoms was thoroughly investigated in a previous theoretical study on 25Na<sub>2</sub>O $xB_2O_3-(75-x)SiO_2$  (x = 0-75 mol %) glasses. That work showed that (i) the positions and areas of the B(III) and B(IV) peaks in the <sup>11</sup>B MAS NMR spectra of the simulated glasses are 965 in good agreement with their experimental counterparts, which 966 validates the accuracy of the structural models and NMR 967 calculations, (ii) the 11B isotropic chemical shift decreases with 968 the B-O-T angle and with the amount of silicon in the second 969 coordination sphere of boron, and (iii) the substitution of 970 B(III)/B(IV) - O - B(III) bonds by B(III)/B(IV) - O - B(IV)971 bonds leads to an increase in the isotropic chemical shift. 972 Unfortunately, the <sup>11</sup>B(III) NMR parameter analyses are not 973 sufficiently distinctive to monitor the changes in the second 974 coordination sphere of these units as P2O5 is added to these 975 glasses. This may be possible, however, by <sup>11</sup>B{<sup>29</sup>Si} REDOR 976 experiments to be conducted on <sup>29</sup>Si enriched glasses in the

Silicon Environment and Connectivity. Table 6 reports the 979 Si<sup>n</sup> distribution in the investigated glasses. In each series, Si is predominantly present as Si<sup>4</sup> species (>84, 78 and 73% for the 981 PB2, MB2, and PA2 series, respectively) and Si<sup>3</sup> species (16, 21, 982 and 26% for the PB2, MB2 and PA2 series, respectively), with small amounts of Si<sup>2</sup> species. While the Si<sup>4</sup> fractions extracted from the MD-generated models are roughly 3-15% higher than 985 those deduced from NMR (Table 4), the trends are the same. 986 With increasing x, the silicate network becomes more 987 polymerized, reflecting the concomitant transfer of the Na<sup>+</sup> ions toward the anionic phosphate component. Table 6 shows 989 that  $Si^4$  is mainly connected to other silica tetrahedrons (1.7–1.8 990 for series PB2, 2.0-2.1 for series MB2, and 2.3-2.4 for series 991 PA2) and B(IV) units (1.5–1.7 for PB2, 1.4–1.6 for MB2, and 992 1.2-1.5 for PA2). Si<sup>3</sup> units, on the other hand, have one NBO 993 which effectively replaces one silicon neighbor in the second 994 coordination sphere. Low amounts of B(III) species are also 995 connected to  $Si^3$  and  $Si^4$  (0.3–0.7). Additionally, there are small 996 fractions of P (up to 0.2 for glasses with larger amounts of  $P_2O_5$ ) 997 that are second neighbors to Si<sup>4</sup> (minimal amounts of P are

Table 6. Amounts of Si<sup>4</sup>, Si<sup>3</sup>, and Si<sup>2</sup> Species (in % of the Si Content) Found in the MD-Generated Glasses and Average Number of NFU Bound to Si<sup>4</sup> and Si<sup>3</sup>  $(\pm 0.1)^a$ 

				(	,	
sample ID	Si <sup>4</sup>	#.	B(IV)	#B(III)	#Si	#P
PB0	84.2		1.7	0.5	1.8	-
PB2-P1	86.4		1.7	0.5	1.7	0.0
PB2-P3	88.5		1.6	0.6	1.7	0.1
PB2-P5	92.3		1.5	0.7	1.7	0.2
PB3-P7	94.8		1.5	0.7	1.5	0.2
MB0	78.8		1.6	0.4	2.1	_
MB2-P1	79.6		1.5	0.4	2.1	0.1
MB2-P3	85.9		1.5	0.5	2.0	0.1
MB2-P5	89.1		1.4	0.4	2.0	0.2
PA0	73.3		1.5	0.3	2.3	_
PA2-P1	83.2		1.2	0.4	2.4	0.0
PA2-P3	80.5		1.3	0.3	2.3	0.1
sample ID	Si <sup>2</sup>	Si <sup>3</sup>	#B(IV)	#B(III)	#Si	#P
PB0	0.1	15.7	1.7	0.4	1.0	_
PB2-P1	0.0	13.7	1.6	0.4	1.0	0.0
PB2-P3	0.1	11.4	1.7	0.3	1.1	0.0
PB2-P5	0.0	7.7	1.5	0.4	1.1	0.0
PB3-P7	0.0	5.3	1.5	0.5	1.0	0.0
MB0	0.3	20.9	1.6	0.3	1.1	_
MB2-P1	0.3	20.1	1.6	0.2	1.2	0.0
MB2-P3	0.0	14.1	1.5	0.3	1.2	0.0
MB2-P5	0.5	10.4	1.4	0.4	1.2	0.1
PA0	1.0	25.7	1.5	0.3	1.2	_
PA2-P1	0.4	16.4	1.2	0.4	1.4	0.0
PA2-P3	0.6	18.9	1.3	0.3	1.3	0.1

 $<sup>^{\</sup>it a}\text{\#}$  indicates the average number of linkages from  $\rm Si^4$  and  $\rm Si^3$  species to each NFU

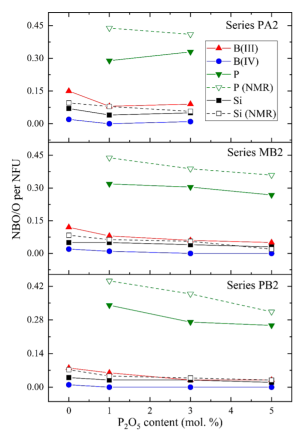
observed around  $\mathrm{Si}^3$  units). As was also observed for B(IV) 998 species, upon addition of  $\mathrm{P_2O_5}$  to the glass, the environment of 999 Si is depleted in B(IV) and Si species and enriched in B(III) 1000 species as the B(III) units tend to link with B(IV) and Si and 1001  $\mathrm{Polester}$  1001

Phosphorus Environment: Short-Range Order. Table 7 1002 17 indicates that the phosphorus speciation in the present glasses is 1003

Table 7. Structural Speciation of Phosphorus (in % of the P Content) in the MD-Generated Glasses

sample ID	P <sup>0</sup> total	P¹ total	P <sup>2</sup> total	P <sup>3</sup> total	P <sup>4</sup> total
PB0	N/A	N/A	N/A	N/A	N/A
PB2-P1	0.0	47.5	41.5	11.0	0.1
PB2-P3	0.0	25.2	58.1	15.3	1.4
PB2-P5	0.4	20.7	60.6	17.2	1.2
PB3-P7	0.0	15.0	73.0	11.2	0.8
MB0	N/A	N/A	N/A	N/A	N/A
MB2-P1	0.0	35.4	56.7	8.0	0.0
MB2-P3	0.9	27.3	64.7	7.2	0.0
MB2-P5	0.0	19.4	68.4	11.8	0.4
PA0	N/A	N/A	N/A	N/A	N/A
PA2-P1	0.1	29.9	55.7	14.3	0.0
PA2-P3	0.9	38.4	52.6	8.3	0.0

dominated by anionic  $P^1$  and  $P^2$  units reflecting the well-known 1004 fact that P preferentially attracts the anionic charges in mixed 1005 network former systems. Quantitative data on this effect are 1006 shown in Figure 9. Plotted are the fractions of NBOs bound to 1007 f9 each NFU in the three glass series studied here, not counting the 1008 uncharged formally doubly bonded NBO already present for any 1009



**Figure 9.** MD-derived average numbers of NBOs per network-forming species B(III), B(IV), Si, and P plotted as a function of x. In the case of the P and Si speciation, results from NMR are shown for comparison.

1010 phosphate unit. The data confirm the results of a previous study 1011 of sodium calcium phospho-borosilicate glasses  $^{38,91}$  indicating 1012 that the attraction of anionic NBO decreases in the order P  $\gg$  1013 B(III) > Si > B(IV). The average number of bridging oxygen 1014 atoms per phosphate unit,  $\langle n \rangle$  as deduced from MD is found to 1015 be somewhat higher than the corresponding experimental values 1016 determined from NMR. Nevertheless, both data sets show that 1017  $\langle n \rangle$  consistently increases with increasing x in the MB2 and PB2 1018 glass series. In contrast, NMR and MD speciation seem to follow 1019 opposite trends in the compositionally more limited series PA2.

*Phosphorus Environment: Connectivity.* Table 8 summa-1021 rizes the NFU connectivities of the phosphate species, indicating 1022 a dominance of Si and B(IV) units. At variance with the MAS 1023 NMR data, the MD results indicate very low levels of P-O-P1024 linkages. One possible reason for this discrepancy can be seen in 1025 the low overall concentration of the P atoms in our glasses. As a 1026 consequence, their speciation and connectivities have little 1027 influence on the energy minimum sought in the simulation. 1028 Considering this situation, the correct prediction of the trend of 1029  $\langle n \rangle$  as a function of  $\alpha$  and the consistent information on P-O-B1030 linkages can already be considered a remarkable success.

Sodium Distribution. The MD simulations reveal that sodium is surrounded by an average of 6.4 and 6.5 oxygen atoms at distances near 2.40 Å in MB0 and PB0 glasses, respectively. Upon addition of  $P_2O_5$  to each glass series, both the average Na coordination numbers and Na-O distances tend to decrease to 6.2/6.3 and 2.37 Å, respectively. The coordination numbers and distances remain almost constant at 6.2 and 2.38 Å, respectively for series PA2. Regarding the partitioning between NBO and BO in the studied glasses, the fraction of NBO around

Table 8. Average Numbers of Network Former Units (B(IV), B(III), P, and Si) Connected to P<sup>1</sup> and P<sup>2</sup> Species in the MD-Generated Glasses<sup>a</sup>

	$\mathbf{P}^1$				
Sample ID	#B(IV)	#B(III)	#Si	#P	
PB2-P1	0.3	0.1	0.6	0.0	
PB2-P3	0.3	0.1	0.6	0.0	
PB2-P5	0.3	0.1	0.5	0.1	
PB3-P7	0.3	0.2	0.3	0.2	
MB2-P1	0.5	0.1	0.4	0.0	
MB2-P3	0.2	0.2	0.6	0.0	
MB2-P5	0.3	0.1	0.6	0.0	
PA2-P1	0.3	0.1	0.6	0.0	
PA2-P3	0.2	0.0	0.7	0.1	
		$P^2$			
sample ID	#B(IV)	#B(III)	#Si	#P	
PB2-P1	1.2	0.0	0.8	0.0	
PB2-P3	1.2	0.2	0.6	0.0	
PB2-P5	0.9	0.2	0.9	0.0	
PB3-P7					
F D3-F /	1.0	0.2	0.7	0.1	
MB2-P1	1.0 1.1	0.2 0.0	0.7 0.9	0.0	
MB2-P1	1.1	0.0	0.9	0.0	
MB2-P1 MB2-P3	1.1	0.0	0.9	0.0	

 $^{\it a}\text{\#}$  indicates the average number of linkages from  $P^1$  and  $P^2$  species to each NFU

Na increases with increasing P<sub>2</sub>O<sub>5</sub> content in the glass, 1040 indicating that sodium gradually shifts from its role as a charge 1041 compensator connected to boron-bonded BO species to a 1042 charge compensator of anionic nonbridging oxygen atoms 1043 attached to phosphate species. To investigate the sodium 1044 distribution around each NFU in the glass, we have plotted the 1045 T-Na (T=Si, B(III), B(IV), P) pair distribution functions in 1046 Figure S6 and compared the T-Na coordination numbers 1047 extracted from the simulations to that computed according to 1048 homogeneously distributed sodium in the glass (see Figure S7). 1049 Figure S6 indicates that the most probable Si-Na distances 1050 range between 3.3 and 3.4 Å and no particular trend is observed 1051 with P2O5 content apart from a slight decrease in their 1052 occurrence (evidenced by a decreased peak intensity). The 1053 B(III)-Na peak is centered at 3.0 Å and splits into two peaks 1054 (3.0 and 3.1 Å) upon P<sub>2</sub>O<sub>5</sub> addition. While the B(III)-Na 1055 interaction is not traditionally expected as these units are charge- 1056 neutral, their spatial proximity can be attributed to either  $B^2(III)$  1057 units containing 1 NBO or B<sup>3</sup>(III) units linked to B(IV) units 1058 which are Na<sup>+</sup> charge-compensated. The B(IV)-Na PDF peak, 1059 on the other hand, is narrower than the B(III)-Na peak and 1060 occurs at slightly shorter distances (2.9 Å), whereas the P-Na 1061 PDFs show a double peak centered at 3.1 and 3.4 Å. From the 1062 viewpoint of the Na+ ions, their interactions with the NFUs in 1063 the second-nearest neighbor sphere are of particular interest. For 1064 defining this sphere, a cutoff distance of 3.6 Å was found to yield 1065 stable results. Table 9 summarizes the development of the Na- 1066 t9 NFU correlations extracted from the MD generated model. 1067 They show a clear dependence on P2O5 content, marked by 1068 monotonic decreases in Si, B(III), and B(IV) neighbors and 1069 increases in the number of P neighbors. Therefore, the sodium 1070 ions strongly prefer phosphate species over any other NFUs in 1071 the second coordination sphere of Na. This finding stands in 1072

1119

Table 9. Contributions of P. B(III), B(IV), and Si to the Second-Nearest Neighbor Coordination of Na in the MD-Generated Glasses

sample ID	#P	#B(III)	#B(IV)	#Si
PB0	-	1.7	2.5	2.3
PB2-P1	0.2	1.6	2.5	2.2
PB2-P3	0.6	1.5	2.2	2.0
PB2-P5	1.0	1.4	2.0	1.7
PB3-P7	1.3	1.3	1.9	1.4
MB0	-	1.4	2.2	2.6
MB2-P1	0.2	1.3	2.2	2.6
MB2-P3	0.7	1.2	1.9	2.3
MB2-P5	1.0	1.1	1.7	2.0
PA0	_	1.0	1.9	3.0
PA2-P1	0.2	0.9	1.7	2.9
PA2-P3	0.6	0.8	1.6	2.5

<sup>&</sup>lt;sup>a</sup># indicates the average number of each NFU within 3.6 Å of Na.

1073 excellent agreement with the conclusions from the <sup>23</sup>Na{<sup>31</sup>P} 1074 REDOR studies. We can put this agreement into more 1075 quantitative terms, considering that the dipolar second moments 1076  $M_{2(S-I)}$ , determined experimentally by REDOR via eq 1 and 1077 listed in Table 3 follow from the internuclear distance 1078 distribution according to the van Vleck equation (eq 7), 92

$$M_{2(S-I)} = \frac{4}{15} \left(\frac{\mu_0}{4\pi}\right)^2 \gamma_I^2 \gamma_S^2 \hbar^2 I(I+1) \sum \frac{1}{r_{S-I}^6}$$
 (7)

1079 where  $\gamma_{\rm I}$  and  $\gamma_{\rm S}$  are the gyromagnetic ratios of the nuclear species 1081 involved, and  $r_{\rm S-I}$  are the internuclear distances between the 1082 observed nuclei (S) and the nuclei (I) of spin quantum number I 1083 to which they are coupled, and all the other symbols have their 1084 usual meanings. As all of the constants in eq 7 are well-known, we can calculate experimental values of  $\sum \frac{1}{r_{s-1}} (NMR)$  from the 1086 numerical second moments  $M_{2(\mathrm{P-Na})}$  and  $M_{2(\mathrm{Na-P})}$  listed in Table 3. This value can be compared with values of  $\sum_{r_{0}} \frac{1}{r_{0}}$ 1088 extracted from the MD generated models, by extending the 1089 summation to the convergence limit, which in the present case 1090 was found to be 15 Å, and results were averaged over a total of 1091 451 <sup>23</sup>Na-observe and 126 <sup>31</sup>P-observe nuclei, respectively. For the glass PB3-P7, these numbers are in very good agreement:

Specifically, we find  $\sum \frac{1}{r_{\rm P-Na}{}^6} = 3.5 \times 10^{57} \, {\rm m}^{-6}$  and  $3.8 \times 10^{57} \, {\rm m}^{-6}$ from NMR and MD output, while the corresponding results for  $\sum \frac{1}{r_{\rm Na-P}} ^{6}$  are  $1.8 \times 10^{57} \, {\rm m}^{-6}$  and  $1.1 \times 10^{57} \, {\rm m}^{-6}$ , respectively. This 1096 preliminary result is very encouraging, showing good consis-1097 tency between the MD and NMR data. Future studies will be 1098 devoted to a more systematic comparison of all the  $M_{2(S-1)}$ 1099 values (S, I =  $^{11}$ B,  $^{23}$ Na,  $^{31}$ P) characterizing all the six possible 1100 pair correlations for a larger set of glasses in the Na<sub>2</sub>O-SiO<sub>2</sub>-1101  $B_2O_3-P_2O_5$  system.

# 5. SUMMARY

1102 The short- and intermediate-range structure of glasses designed 1103 in the peralkaline, metaboric, and perboric regions of the Na<sub>2</sub>O-1104 B<sub>2</sub>O<sub>3</sub>-P<sub>2</sub>O<sub>5</sub>-SiO<sub>2</sub> system has been investigated. In general, it is 1105 observed that an increase in the P2O5 content in the glasses 1106 produces an increase in the phosphate connectivity in the 1107 network ( $P^2/P^3$  units replacing isolated  $P^0/P^1$  units). This is 1108 accompanied by a restructuring of the borate network (favoring B<sup>3</sup>(III) species) and a considerable repolymerization of the 1109 silicate network. These changes are attributed to a significant 1110 redistribution of Na+ cations toward anionic phosphate NBO 1111 sites and away from silicate or borate network units. The rate of 1112 change in average P connectivity to boron  $\langle n_B(P) \rangle$  as a function 1113 of  $P_2O_5$  content increases in the order of PA  $\rightarrow$  MB  $\rightarrow$  PB. The 1114 present study demonstrates that MD simulations can be used to 1115 guide and/or validate the structural interpretation of MAS NMR 1116 data. In addition, MD simulations can provide useful 1117 supplemental structural data not available from NMR studies. 1118

## 6. IMPLICATIONS ON THE DESIGN OF NOVEL **BOROSILICATE BIOACTIVE GLASSES**

It is well-known that the short- and intermediate-range order in 1120 multicomponent glasses greatly affects the glass degradation 1121 behavior in simulated body environments. For instance, weakly 1122 bound species (i.e., Na+ at Si-NBO sites) are readily released 1123 from the glass in aqueous environments via ion exchange. 93-95 Also, B(III) species (acting as Lewis acid sites) have been 1125 previously shown to react with water at quicker rates in 1126 comparison to B(IV) species. Furthermore, isolated 1127 orthophosphate (P0) units are known to be released easily 1128 from the glass and to promote biomineralization, whereas  $P_{Si}^{1}$  1129 units inhibit the same bioactive processes. However, in 1130 the present phospho-borosilicate glass system, significant 1131 amounts of weakly bound isolated phosphate species (P<sup>0</sup> and 1132  $P_{1p}^{1}$ ) in the x = 1 glasses are replaced by species containing P- 1133 O-B(IV) NFU linkages. The latter species, similarly as  $P_{1S_i}^1$  1134 units, are likely to improve the chemical durability of glass, thus, 1135 adversely impacting their bioactive response. Furthermore, the 1136 additional structural effects that P2O5 exhibits upon the NFU 1137 mixing and speciation (i.e., B(IV) and Si<sup>4</sup> fractions), including 1138 its impact on the distribution and proximity of Na<sup>+</sup> to specific 1139 anionic sites, can have profound effects on the ionic release 1140 characteristics and bioactivity which can benefit the conception 1141 of novel bioactive glasses for particular applications. Future 1142 studies examining structure-degradation behavior relationships in 1143 the present sodium phospho-borosilicate glass system can help 1144 to uncover the ideal NFU structures necessary for tuning ionic 1145 release to elicit desired biological responses when in contact 1146 with human body fluids. The described approach, which 1147 necessitates a careful compositional design, can accelerate the 1148 development of borosilicate glasses for innovative biomedical 1149 applications.

# 7. CONCLUSIONS

The present study combines the strengths of state-of-the-art 1151 experimental and computational techniques to provide a 1152 comprehensive structural understanding of the short- and 1153 intermediate-range ordering in the Na<sub>2</sub>O-P<sub>2</sub>O<sub>5</sub>-B<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> 1154 based model bioactive glass system. The impact of P<sub>2</sub>O<sub>5</sub> on the 1155 structure of glasses designed in the perboric (R < 1), metaboric 1156 (R = 1), and peralkaline (R > 1) regions of the above-mentioned 1157 system has been investigated. While  $P_2O_5 > 3$  mol % leads to 1158 phase separation and formation of crystalline Na<sub>4</sub>P<sub>2</sub>O<sub>7</sub> in 1159 peralkaline systems, the glass-forming ability increases with 1160 decreasing R-value. Furthermore, P2O5 induces repolymeriza- 1161 tion in the silicate units and a restructuring of the borate 1162 component in the glass network, which is driven by the 1163 additional demand of phosphate for charge compensation. In 1164 addition, the degree  $\langle n \rangle$  of phosphate polymerization increases 1165 as a function of P<sub>2</sub>O<sub>5</sub> content, as increasing amounts of B-O-P 1166

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1167 linkages are formed. The latter also serve to disperse the negative 1168 charges of the B(IV) units toward nonbridging oxygen atoms 1169 located on the phosphate species.  $^{87,88}$ 

The detailed analysis of <sup>31</sup>P-<sup>11</sup>B dipole-dipole and <sup>31</sup>P-<sup>31</sup>P 1171 indirect spin-spin couplings on selected samples suggests a 1172 general picture of the NFU connectivity, which stands in 1173 excellent agreement with the conclusions drawn from the MD 1174 simulations and expands significantly on previous findings on bioactive P<sub>2</sub>O<sub>5</sub>-containing silicate and borosilicate glasses. <sup>30,38,91</sup> 1176 In addition, the combined <sup>23</sup>Na-based double resonance experiments and the MD simulations portray a consistent and quantitative picture of the next-nearest neighbor coordination 1179 sphere of the modifier cations, featuring their preferred 1180 correlations with the phosphate component. Ultimately, the comprehensive structural details presented here will allow us to 1182 develop predictive models for dissolution kinetics and 1183 bioactivity. This will facilitate the compositional design of 1184 gene-activating borosilicate-based glasses with release kinetics 1185 that can be tailored to the specific demands of individual 1186 therapeutic applications.

#### 187 ASSOCIATED CONTENT

# 1188 Supporting Information

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The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.0c04470.

XRD patterns of the studied glasses; supplemental single resonance MAS NMR data, PDF data for the MD-simulated glasses, Na-NFU coordination environment extracted from MD simulations, experimental parameters used for single and double resonance MAS NMR experiments, number of atoms utilized for each MD-simulated glass, interatomic potential parameters used for MD simulations, and fitting parameters and calculated fraction of each species for all <sup>11</sup>B and <sup>31</sup>P MAS NMR data (PDF)

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