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Introduction

Narrow-emission and highly efficient blue thermally activated delayed fluorescence (TADF) materials approaching the Rec.2020 standard for color gamut (CIE coordinates: (0.131, 0.046), as recommended by the International Telecommunication Union) have attracted significant research attention due to their immense potential in the field of 4K/8K ultrahigh-definition displays.^{1,2} However, the development of such materials has encountered considerable challenges, as it requires concurrent optimization of the high photoluminescence (PL) efficiency (Φ_{PL}), narrow full width at half maximum (FWHM), fast reverse intersystem crossing (RISC) process, and wide bandgap. Therefore, only a handful of reports exist on blue TADF organic light-emitting diodes (OLEDs) with a CIE_y value

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Realizing highly efficient deep-blue organic lightemitting diodes towards Rec.2020 chromaticity by restricting the vibration of the molecular framework[†]

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Deep-blue organic light-emitting diodes (OLEDs) with narrow emission spectra and high efficiency, meeting the Rec.2020 standard, hold significant promise in the realm of 4K/8K ultrahigh-definition displays. However, the development of light-emitting materials exhibiting both narrowband emission and high efficiency, particularly in the realm of deep-blue thermally activated delayed fluorescence (TADF), confronts substantial challenges. Herein, a novel deep-blue TADF emitter, named **BOC-PSi**, was designed by integrating a rigid B-heterotriangulene acceptor (A) with a rigid phenazasiline donor (D). The replacement of a sp³ carbon atom with a sp³ silicon atom in the D moiety helps to restrict the low-frequency bending vibration throughout the entire D-A molecular backbone, while concurrently accelerating the multi-channel reverse intersystem crossing (RISC) processes. Notably, OLEDs using the **BOC-PSi** emitter exhibit exceptional performance, with a high maximum external quantum efficiency (EQE_{max}) approaching 20%, and a superior color purity closely aligning with the Rec.2020 blue standard.

close to 0.046 and a maximum external quantum efficiency (EQE_{max}) surpassing 15%.³⁻⁷

To achieve TADF materials with a narrow emission bandwidth, it is essential to impart a rigid framework to the fluorophore. A notable example of this concept is multiple resonance (MR) B,N-heteroarenes, known for exhibiting narrow FWHM and high $\Phi_{\rm PL}$ due to their highly rigid molecular scaffolds.⁸⁻¹⁰ Nevertheless, their robust planar molecular backbones significantly hinder the realization of strong spin-orbit coupling (SOC).¹¹ As a result, most MR-TADF dyes experience relatively slow RISC processes, posing challenges for their implementation in conventional TADF-OLEDs without the need supplementary TADF sensitizers.

On the other hand, due to the compensating effect between their variations in orbital angular momentum and spin angular momentum,^{12,13} TADF molecules with highly distorted donoracceptor (D–A) structures are more likely to exhibit a relatively fast RISC process. However, as the D and A moieties are chemically linked only through a fragile single bond, these D–A dyads generally suffer from poor framework rigidity, leading to a relatively wide FWHM and an accelerated non-radiative process. Moreover, to minimize the singlet–triplet energy gap ($\Delta E_{\rm ST}$), these D–A TADF compounds often possess a nearly orthogonal orientation between their D and A units, resulting in a reduced radiative transition rate. Consequently, the accelerated non-radiative process poses a significant obstacle to achieving a high $\Phi_{\rm PL}$.



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Fig. 1 (a) Molecular and single crystal structures of BOC-PC and BOC-PSi; (b) PL spectra of PC, PSi, BOC-PC and BOC-PSi (in toluene at a concentration of 10 μ M); (c) PL decay curves of BOC-PC- and BOC-PSi-based film samples (15 wt% in DPEPO under N₂); (d) calculated Huang–Rhys factors and corresponding vibration modes contributed significantly to the reorganization energy of BOC-PC and BOC-PSi from the S₁ to S₀ states (PBE0/6-31G** at optimized S₁ geometry).

In addition, to obtain TADF D–A dyads with deep-blue emission performance approaching the Rec.2020 standard, it is necessary to carefully select D and A moieties not only having a deep highest occupied molecular orbital (HOMO) and a shallow lowest unoccupied molecular orbital (LUMO), respectively, but also possessing a local triplet excited state (³LE) energy level higher than 3.0 eV. However, such structural units are scarce, which further adds to the challenge of constructing TADF emitters with a wide bandgap.¹⁴

Herein, we report a high-performance deep-blue TADF D–A dyad, namely **BOC-PSi**, which utilizes a rigid B-heterotriangulene derivative (BOC) as the A moiety, and 10,10-diphenyl-5,10-dihydrodibenzo[*b*,*e*][1,4]azasiline (PSi), the sila-product of 9,9-diphenyl-9,10-dihydroacridine (PC), as the D moiety (Fig. 1a). **BOC-PSi**-based OLEDs demonstrated an EQE_{max} of 19.6% under a CIE_y value of 0.049, establishing **BOC-PSi** as an advanced Rec.2020 blue OLED emitter (Table S3†). Comparative studies between **BOC-PSi** and **BOC-PC** (a reference compound using PC as the D unit) revealed that the better electroluminescence (EL) performance of **BOC-PSi** can be mainly attributed to its narrower FWHM and higher Φ_{PL} , which arise from the enhanced rigidity of PSi compared to PC. These findings demonstrated that D–A TADF dyads with exceptional blue color gamut can be acquired by employing rigid D and A subunits.

Molecular design rationale & characterization

The design rationale for **BOC-PSi** is as follows. (1) To endow the dyad with a highly twisted geometry, PC composed of merely

six-membered polycyclic ring systems was chosen as the parent D scaffold.¹⁵ (2) A sila-modification was performed on PC, because the resulting PSi shows better framework rigidity compared to PC, as evidenced by its narrower and more structured fluorescence and phosphorescence spectra (Fig. 1b and S8†). (3) The ³LE energy level of PSi is as high as 3.10 eV, and its calculated HOMO (-5.60 eV) is deeper than that of PC, which may be attributed to the weaker hyperconjugation effects in PSi (Fig. S11†).^{16,17} These conditions facilitate the acquisition of a TADF D-A dyad with a wide bandgap. (4) Based on the HOMO energy level data of PSi, BOC was screened out as the A moiety due to its integrated rigid molecular skeleton, high ³LE energy level of 3.17 eV, and shallow LUMO level of -2.36 eV.¹⁸

The synthetic routes of **BOC-PSi** and **BOC-PC** are illustrated in the ESI (Scheme S1).† The molecular structure of **BOC-PSi** and **BOC-PC** was confirmed by nuclear magnetic resonance spectroscopy (NMR), high-resolution mass spectroscopy (MS), and single crystal X-ray diffraction (XRD).

The single crystal structures of **BOC-PSi** and **BOC-PC** are depicted in Fig. S10,† and the corresponding crystal parameters are listed in Table S1.† As expected, **BOC-PSi** and **BOC-PC** both show a nearly perpendicular orientation between their D and A segments, with torsion angles measuring 86.4° and 84.0°, respectively. This highly twisted geometry offers advantages in promoting efficient RISC by facilitating enhanced vibronic coupling between the charge-transfer triplet excited state (³CT) and ³LE state. However, the two phenyl substituents of D units in the two compounds have quite different relative positions. As shown in Fig. 1a, the two phenyls of **BOC-PSi** are nearly

Chemical Science

symmetrically distributed above and below the azasiline plane, whereas those of **BOC-PC** exhibit distinct conformations, with one quasi-axial and the other quasi-equatorial. The disparity in the relative positions of the phenyls of **BOC-PS** and **BOC-PC** can be attributed to the more planar configuration of the azasiline ring compared to the acridine ring (dihedral angle: ~6° vs. ~24°, Fig. 1a and S10†), which is due to the significantly longer C_{sp^2} -Si_{sp³} bonds than the corresponding C_{sp^2} -C_{sp³} bonds (~1.8 vs. ~1.5 Å, Table S1†), resulting from the larger atomic radius of silicon than carbon.

Electrochemical & thermal stability properties

Through cyclic voltammetry (CV) measurements (Fig. S5†), the HOMO energy level values of **BOC-PC** and **BOC-PSi** were determined to be -5.41 eV and -5.53 eV, respectively. The deeper HOMO level of **BOC-PSi** than **BOC-PC** indicates the weaker electron-donating ability of PSi compared to PC, which ultimately benefits the realization of wider bandgap emission. Subsequently, the LUMO energy level values were calculated from the corresponding optical bandgap and HOMO energy level data, and were determined to be -2.32 eV and -2.37 eV for **BOC-PC** and **BOC-PSi**, respectively. The slight disparity in the LUMO energy level values between the two compounds could potentially stem from the variation in the D–A dihedral angles in their optimized S₀ geometrical structures (Fig. S12†).

Based on the results of thermal gravimetry analysis (TGA) and differential scanning calorimetry (DSC) characterizations (Fig. S6†), both compounds demonstrate excellent thermal stability, as evidenced by their high decomposition temperatures (T_d) exceeding 340 °C at 5% initial weight loss. In addition, only **BOC-PC** exhibits a notably high melt temperature (T_m) of 327 °C, while no significant endothermic signal ascribed to glass transition could be observed in both emitters.

Photoluminescence properties

The photophysical properties of these two emitters were investigated in dilute solution (10^{-5} M) and doped film states (15 wt% in bis[2-(diphenylphosphino)phenyl]ether oxide (DPEPO)). As illustrated in Fig. 1b and Table 1, **BOC-PSi** and **BOC-PC** both emit deep-blue PL in toluene, but **BOC-PSi** exhibits a superior blue color gamut to **BOC-PC** due to its >10 nm blue-shifted PL emission band (λ_{PL} : 432 vs. 445 nm) and much narrowed FWHM (61 vs. 70 nm). These findings provide clear evidence for the effectiveness of this strategy in the rational design of deep-blue emitters. Besides, with increasing solvent polarity from hexane to acetonitrile, both **BOC-PSi** and **BOC-PC** display red-shifted and broadened PL spectra, manifesting the CT character of their S₁ states (Fig. S7†). Notably, **BOC-PSi** consistently shows a narrower PL spectrum compared to **BOC-PC** in every solvent, suggesting that a more rigid D subunit can indeed induce a narrower FWHM of the corresponding ¹CT-featured emission in a D-A dyad.

In terms of the two film samples, they both exhibit slightly narrowed PL spectra compared to their corresponding toluene solutions (FWHM: 55 vs. 61 nm for **BOC-PSi**; 66 vs. 70 nm for **BOC-PC**), which can be attributed to the restricted rotation of the C–N single bond within a more rigid matrix. In comparison to **BOC-PC**, **BOC-PSi** also shows a blue-shifted (λ_{PL} : 439 vs. 454 nm) PL spectrum with a narrowed FWHM (55 vs. 66 nm). Additionally, **BOC-PSi** shows a higher Φ_{PL} compared to **BOC-PC** (92% vs. 80%), indicative of the existence of additional exciton loss pathways in **BOC-PC**, potentially arising from the vibration relaxation of the fluorophore.

Further transient PL measurements revealed the presence of delayed fluorescence (DF) behavior in both film samples. Notably, the DF lifetime (τ_{DF}) is as short as 2.6 µs for **BOC-PSi** and 1.8 µs for **BOC-PC** (Fig. 1c), indicative of the occurrence of fast RISC processes in both **BOC-PSi** and **BOC-PC**. With respect to the prompt fluorescence (PF), the average lifetime (τ_{PF}) is determined to be 9.4 ns for **BOC-PSi** and 14.2 ns for **BOC-PC** (Fig. S9†). Based on the τ_{DF} , τ_{PF} , Φ_{PF} and Φ_{DF} data (Table 1), the rate constants for key photophysical processes, including fluorescence decay (k_F), intersystem crossing (k_{ISC}), RISC (k_{RISC}), and non-radiative decay of the S₁ state (k_{NR} ^S), were calculated for the two compounds.^{19,20}

Exciton dynamics process

As shown in Table 1, **BOC-PSi** has a significantly smaller $k_{\rm NR}^{\rm S}$ than **BOC-PC** $(2.41 \times 10^6 \text{ vs. } 8.63 \times 10^6 \text{ s}^{-1})$. To elucidate the underlying cause of the suppressed non-radiative process resulting from the sila-modification, theoretical calculations were conducted to determine the total reorganization energy (λ_{total}) between the S₁ and S₀ states for both compounds as well as the A moiety of BOC. The results indicated that the λ_{total} of BOC-PSi is slightly smaller than that of BOC-PC (0.20 vs. 0.30 eV, vide Fig. 1d), implying that BOC-PSi possesses superior molecular skeleton rigidity to **BOC-PC**,^{21,22} while the λ_{total} of BOC is only 0.17 eV, indicative of its excellent rigidity (Fig. S14⁺). In line with this deduction, the root mean square deviation (RMSD) value of the superposition of the optimized S₀ and S₁ geometries of BOC-PSi is much smaller than that of BOC-PC (0.123 vs. 0.316 Å, Fig. S15[†]).²⁴ These findings suggest a stronger suppression of non-radiative decay for the S₁ state of BOC-PSi than BOC-PC.

Table 1Key photophysical data of BOC-PC and BOC-PSi (15 wt%-doped in DPEPO under N_2)												
Compound	$\lambda_{\mathrm{PL}}\left[\mathrm{nm} ight]$	$\Phi_{ m PL}$	$\Phi_{ m PF}$	$\Phi_{ m DF}$	$\Phi_{ m ISC}$	$\tau_{\rm PF} [\rm ns]$	$\tau_{\rm DF}\left[\mu s\right]$	$k_{ m F} \left[{ m s}^{-1} ight]$	$k_{\rm ISC} [{ m s}^{-1}]$	$k_{ m RISC} [m s^{-1}]$	$k_{\rm NR}{}^{\rm S}\left[{ m s}^{-1} ight]$	
BOC-PC BOC-PSi	454 439	80% 92%	49% 26%	31% 66%	39% 72%	14.2 9.4	1.8 2.6	$3.45 imes10^7\ 2.77 imes10^7$	$2.73 imes10^7$ $7.63 imes10^7$	$\begin{array}{c} 9.07\times10^5\\ 1.36\times10^6\end{array}$	$\begin{array}{c} 8.63\times10^6\\ 2.41\times10^6\end{array}$	

To elucidate the reason for the disparity in FWHM between the two emitters, the Huang-Rhys factors (HRf) at various vibration modes were calculated for BOC-PSi, BOC-PC and BOC. For BOC-PC, a low-frequency scissoring swing of the entire molecular framework was observed at a normal mode wavenumber of 10 cm^{-1} (Fig. 1d and S13[†]), accompanied by a large HRf of 4.25.23 Detailed vibration mode analysis revealed that the scissoring motion along the D-A skeleton in BOC-PC can be ascribed to the top-heavy nature of its PC moiety during the bending vibration of the C-N bond. Additionally, the vibration mode in BOC-PC that exhibits the second-largest HRf (0.82) also arises from the twisting of the PC moiety. In contrast, due to the well-balanced character of its PSi moiety, all HRfs calculated at the low-frequency region below 200 cm⁻¹ are significantly smaller than those of BOC-PC, and no obvious vibrational motions throughout the whole D-A scaffold of BOC-PSi were observed. Therefore, the BOC-PSi exhibits a smaller overall HRf compared to BOC-PC, manifesting a significantly suppressed structural relaxation thus narrowing the FWHM.²⁴ In the case of BOC, no detectable vibrations were found contributing to its HRf in the low-frequency range below 200 cm⁻¹, indicative of its excellent skeletal rigidity. Therefore, it can be inferred that the severe non-radiative process and larger FWHM in BOC-PC should be mainly ascribed to its PC subunit.

Excitingly, the $k_{\rm ISC}$ and $k_{\rm RISC}$ values of **BOC-PSi** are also both larger than those of **BOC-PC** ($k_{\rm ISC}$: 7.63 × 10⁷ vs. 2.73 × 10⁷ s⁻¹; k_{RISC} : 1.36 × 10⁶ vs. 9.07 × 10⁵ s⁻¹), indicating a stronger SOC effect and/or a smaller energy difference between the S_1 and $T_1/$ T_n states in **BOC-PSi**. To understand the reason behind the larger k_{RISC} value of **BOC-PSi** than **BOC-PC**, the PL and phosphorescence (Phos) spectra of both compounds were recorded at 77 K. The structureless and red-shifted PL and Phos spectra of the two compounds relative to their corresponding D/A fragments manifest the ¹CT and ³CT features of their S₁ and T₁ states, respectively (Fig. S8[†]). The ¹CT/³CT energy levels, according to the onset of the PL and Phos spectra, were estimated to be 3.06/2.99 eV for BOC-PC and 3.11/3.05 eV for BOC-PSi. Although the singlet-triplet splitting of the CT excited states in BOC-PC and BOC-PSi is guite similar (0.07 vs. 0.06 eV), there is an evident difference in the energy splitting between their ¹CT and ${}^{3}LE_{D}/{}^{3}LE_{A}$ states. As depicted in Fig. S8,[†] the ${}^{3}LE$ energy levels of PC and PSi were both calculated to be approximately 3.20 eV, while that of BOC was estimated to be 3.35 eV. Therefore, the absolute values of $\Delta E_{\rm ST}$ (¹CT-³LE_D) and $\Delta E_{\rm ST}$ (¹CT-³LE_A) of BOC-PC are 0.14 eV and 0.29 eV, respectively, whereas those for BOC-PSi are 0.09 eV and 0.24 eV respectively. Considering that D-A dyads with highly twisted molecular geometries typically exhibit a much stronger SOC effect between a ³LE and a ¹CT state compared to that between a ³CT and a ¹CT



Fig. 2 (a) Calculated excited states energy and NTO distributions for BOC-PC and BOC-PSi; (b) the plausible mechanism of harvesting triplet states for BOC-PC (top) and BOC-PSi (bottom); (c) EQE as a function of the current density (*J*) of BOC-PC-based device I, and BOC-PSi-based device II (inset: EL spectra of devices I–II at *J* of 10 mA cm⁻²); (d) current density–voltage–luminance (J-V-L) profiles of devices I–II; (e) comparison of EQE_{max} for the reported deep-blue OLED with CIE_v of 0.03–0.06.

Table 2 The key EL properties of BOC-PSi- and BOC-PC-based OLEDs

Dopant	Device	$\lambda_{ m EL}$ (nm)	FWHM (nm)	CE_{max} (cd A^{-1})	EQE^{a}/EQE^{b} (%)	Roll-off ^c	$CIE_{1931}(x, y)$
BOC-PC	Ι	450	62	11.12	14.8/2.8	85%	(0.148, 0.085)
BOC-PSi	II	433	53	9.10	19.6/14.8	24%	(0.154, 0.049)
^a Device max	ximum external	l quantum effi	ciency. ^b External qua	antum efficiency at 100	cd m ⁻² . ^c Efficiency roll	off from 1 cd m⁻	$^{-2}$ to 100 cd m $^{-2}$.

state,¹³ the faster RISC process in **BOC-PSi**, as compared to **BOC-PC**, may be attributed to the smaller absolute values of $\Delta E_{\rm ST}$ (¹CT-³LE).

This deduction was supported by theoretical computations. As depicted in Fig. 2a, the S₁ and T₁ states of BOC-PC and BOC-PSi were both calculated to show a CT feature, and the energy splitting between the two states is 0.01 eV for both compounds. In line with our conjecture, the calculated SOC constants for the $T_1 \rightarrow S_1$ process are 0.04 cm⁻¹ for **BOC-PC** and 0.03 cm⁻¹ for BOC-PSi, both are too small to trigger fast RISC processes. Nevertheless, the T₂ states of BOC-PC and BOC-PSi are both dominated by the BOC unit, displaying a ${}^{3}LE_{A}$ character. Despite having an identical T_2 energy level (~3.0 eV), the lower S_1 state of **BOC-PC** results in a larger absolute value of ΔE_{ST} $(^{1}CT-^{3}LE_{A})$ compared to **BOC-PSi** (0.18 vs. 0.06 eV). Considering that the SOC constants between the T₂ and S₁ states of BOC-PC and **BOC-PSi** are relatively large (>0.70 cm⁻¹), both compounds are expected to undergo a relatively fast $T_2 \rightarrow S_1$ RISC process. However, **BOC-PSi** is likely to achieve a larger k_{RISC} due to its much smaller $\Delta E_{\rm ST}$ (¹CT-³LE_A) value compared to BOC-PC. Additionally, for BOC-PSi, its T₃ state (exhibiting a ³LE_D character) was found to be close to its S₁ state ($\Delta E_{ST} = 0.20$ eV), and the calculated SOC constant for the $T_3 \rightarrow S_1$ process was also substantial at 0.58 cm⁻¹, implying the presence of a fast $T_3 \rightarrow$ S_1 RISC process in **BOC-PSi**. Consequently, the large k_{RISC} of BOC-PSi is believed to stem from its effective multi-channel RISC processes.

Therefore, through the substitution of the sp³-C atom within the 9,10-diphenylacridine segment of **BOC-PC** with a sp³-Si, we have acquired **BOC-PSi**, which has a better-balanced and robust molecular framework, a slightly deepened HOMO energy level, and a maintained high ³LE energy level. As a result, in comparison to **BOC-PC**, **BOC-PSi** shows a wider emission bandgap, a narrower FWHM, a more suppressed non-radiative process and hence a higher Φ_{PL} , as well as a faster RISC process. Consequently, it is expected that **BOC-PSi** will demonstrate superior EL performance to **BOC-PC**.

Electroluminescence performance

Subsequently, OLEDs were fabricated using **BOC-PSi** or **BOC-PC** as the doping guest (15 wt% in DPEPO): ITO/PEDOT: PSS (30 nm)/TAPC (30 nm)/TCTA (10 nm)/mCP (10 nm)/DPEPO: emitters (15 wt%, 40 nm)/DPEPO (5 nm)/TmPyPB (35 nm)/LiF (1.2 nm)/Al (120 nm). Consistent with our conjecture, the **BOC-PSi**-based device II shows significantly superior EL performance to the **BOC-PC**-based device I. As depicted in Table 2, device I

displayed an inferior EL color purity than device II (CIE_y: 0.085 vs. 0.049) due to the red-shifted EL band (λ_{EL} : 450 vs. 433 nm) and wide FWHM (62 vs. 52 nm). Besides, device I also exhibited a lower EQE_{max} (14.8% vs. 19.6%) together with more pronounced efficiency roll-off. These disadvantages can be ascribed to the narrower emission bandgap, wider FWHM, lower Φ_{PL} and slower RISC process of **BOC-PC** than **BOC-PSi**. All these findings confirm the potential of **BOC-PSi** as a more promising deep-blue OLED emitter than **BOC-PC**. It is noteworthy that the **BOC-PSi**-based OLEDs stand as one of the top-performing examples among reported deep blue TADF OLEDs whose color purity approaches the Rec.2020 blue standard.

Conclusion

In conclusion, we demonstrated that by employing rigid D and A structural units, TADF D-A dyads with narrow FWHM and high $\Phi_{\rm PL}$ can be developed. Using **BOC-PSi** as an example, we proved that 10,10-diphenyl-5,10-dihydrodibenzo[b,e][1,4]azasiline (PSi) is a more promising D unit than 9,9-diphenyl-9,10dihydroacridine (PC) when constructing deep blue TADF D-A dyads with good color gamut. The reason for this is that the replacement of a sp³-C with a sp³-Si helps to restrict the lowfrequency bending vibration along the whole D-A molecular backbone, thus minimizing exciton energy loss, while concurrently accelerating the multi-channel RISC processes. OLEDs using BOC-PSi as the emitter exhibited not only an impressive EQE_{max} of nearly 20% and a narrow FWHM of 53 nm, but also a superior color purity approaching the Rec.2020 blue standard. This work would provide a new avenue in developing highly efficient, narrow-emission and wide-bandgap blue TADF-OLED materials.

Data availability

Experimental procedures, details of the calculations, and additional data can be found in the ESI.†

Author contributions

C. L., K. Z. and Y. L. contributed equally to this work. C. L. – conceptualization, synthesis, investigation, visualization, writing – original draft; K. Z. – OLED devices, investigation; Y. L. – investigation, funding acquisition, writing – review & editing; Y. Y. – investigation (photophysics); Y. H. – investigation (photophysics); M. J. – investigation (photophysics); Y. H. – investigation (photophysics); Y. L. – synthesis; J. T. – funding acquisition, OLED devices, review & editing; Y. H. – review & editing; Z. L. – funding acquisition, project administration, writing – review & editing, conceptualization, supervision.

Conflicts of interest

There are no conflicts to declare.

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