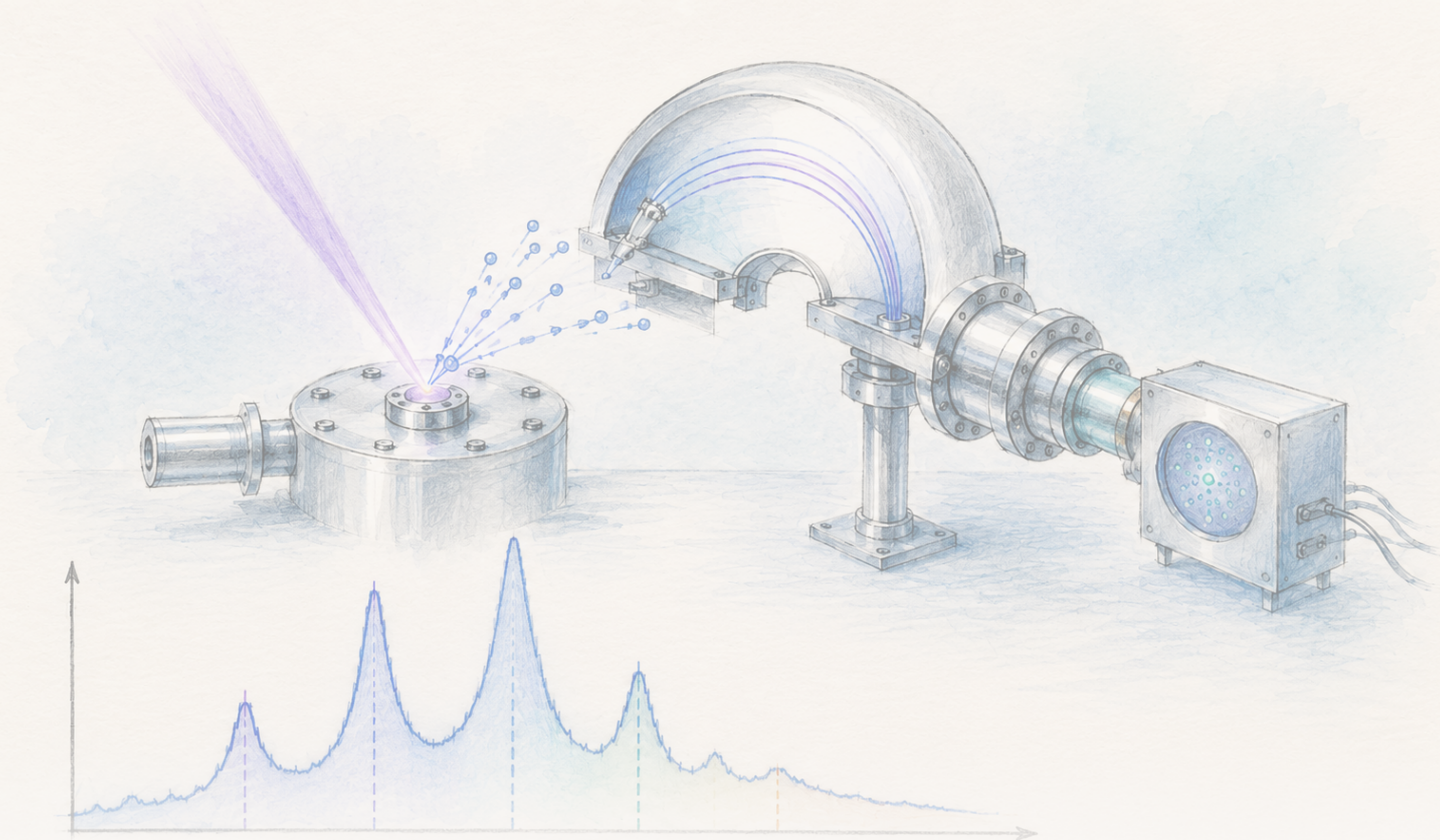




The CLEAN PAPER

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Carbon-bismuth triple bond inaonyesha wapi textbook sigma-na-pi picture inaacha kufanya kazi

Utafiti wa Science unaprobe CBi- molecular ion, heavy-element cousin wa familiar triply bonded species, kwa cryogenic photoelectron spectroscopy na relativistic quantum calculations. Result ni direct evidence kwamba strong spin-orbit coupling inaweza kureorganize classical sigma na pi bonding framework kuwa relativistic Kramers pairs zilizo labeliwa kwa total angular momentum. Hii haifanyi ordinary chemical bonding kuwa wrong; inaonyesha kwa nini bookkeeping inabadi- lika karibu na very heavy atoms.

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Paper: *Relativistic collapse of the classical triple bond in the CBi- molecular ion*, Deniz Kahraman, Jie Hui, Xin-Yu Zhang, Neil A. Ellis, Hyun Wook Choi, Kirk A. Peterson and Lai-Sheng Wang, Science 393, 184-187 (2026) · DOI: 10.1126/science.aei1285

Triple bond kwa kawaida ni safi. Bismuth inaifanya iwe na fujo zaidi.

Picha ya kawaida ya triple bond ni sigma bond moja na pi bonds mbili. Sigma bond ni sehemu ya bond inayokutana head-on, ikiwa na electron density kwenye mstari kati ya atoms mbili. Pi bond ni sehemu ya side-on, ikiwa na electron density juu na chini, au kuzunguka, mstari huo. Katika textbook triple bond, sigma moja pamoja na pi mbili hutengeneza package safi.

Ni mchoro safi, na kwa light atoms mara nyingi ni safi kwa sababu nzuri: electrons zinaweza kuelezwa katika orbitals ambazo shapes na spins zake zinaweza kutenganishwa kimawazo.

Picha hiyo si uongo. Ni approximation nzuri sana katika sehemu ya periodic table ambako textbook examples nyingi zinaishi.

Bismuth haiko katika sehemu hiyo ya periodic table.

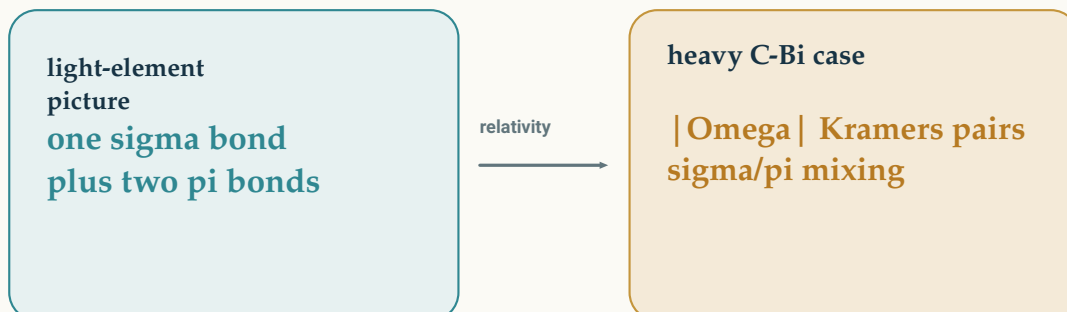
Bismuth ina protons 83. Karibu na nucleus nzito kiasi hicho, relativity huacha kuwa correction ya mapambo na kuwa sehemu ya chemistry. Electrons husogea katika field yenye nguvu kiasi kwamba spin-orbit coupling — coupling kati ya spin ya electron na orbital motion yake — inaweza kuwa moja ya facts kuu zinazopanga mfumo. Hilo likitokea, old labels hazipotei kwa sababu mtu amezisahau. Zinaacha kuwa labels bora.

makala mpya ya Science ya Deniz Kahraman, Jie Hui, Xin-Yu Zhang, Neil A. Ellis, Hyun Wook Choi, Kirk A. Peterson na Lai-Sheng Wang inasoma case iliyochaguliwa kwa ukali: carbon-bismuth molecular ion CBi⁻. Ni isovalent na CN⁻, light-element triple-bond mfumo inayojulikana. Lakini kubadilisha nitrogen na bismuth kunahamisha electron-counting problem ileile katika relativistic environment nzito zaidi sana.

matokeo safi si “triple bonds ni wrong.” Ni finyu na ya kuvutia zaidi: katika CBi⁻, classical sigma-plus-two-pi description inaporomoka na kuwa relativistic description iliyojengwa kwa Kramers pairs zilizo labeliwa kwa total angular-momentum projection. Bond bado ipo. Bookkeeping ya zamani ndiyo inayoshindwa.

The textbook triple bond meets heavy-atom relativity

Spin-orbit coupling mixes the clean sigma/pi picture in C-Bi.



Boundary: the textbook model still works where relativity is small.

Light-element triple bond ni sigma orbital moja plus pi mbili; katika heavy C-Bi, spin-orbit coupling inapanga upya mfumo kuwa $|\Omega|$ Kramers pairs zenye sigma/pi mixing. Bond bado ipo — textbook labels ndizo zinaacha kufanya kazi vizuri.

Original diagram — The Clean Paper CC BY 4.0

Waandishi walipima nini

Waandishi walitengeneza CBi- katika molecular beam kwa laser ablation ya bismuth/graphite target, kisha wakaiprobe anion kwa high-resolution cryogenic photoelectron spectroscopy na photoelectron upigaji picha. Anion ni ion yenye charge hasi; hapa CBi- ni carbon-bismuth molecule yenye electron moja ya ziada.

Photoelectron spectroscopy hufanya jambo rahisi kwa njia technical. Photon huondoa electron kutoka anion. Kupima outgoing electron hukuambia energy kiasi gani kilihitajika, na hivyo neutral electronic state ipi ilifikiwa. Neutral electronic state ni arrangement moja inayoruhusiwa ya electrons zilizobaki baada ya extra electron kuondolewa. Photoelectron upigaji picha huongeza angular information: inarekodi pattern ya emitted electrons, inayosaidia kutambua character ya orbital electron ilikotoka.

[video pending]

Educational animation fupi ya photoemission athari: incoming light huondoa electrons kutoka material, na energies za electrons zinaonyesha states zilizoachwa nyuma — principle ileile nyuma ya photoelectron

spectroscopy inayotumika hapa. Inaonyesha general technique, si CBI- experiment yenyewe. Imetranscodeiwa kuwa MP4 na The Clean makala kwa browser compatibility; content haijabadilishwa.

Spectrum kuu katika 4.661 eV ilionyesha bendi tatu za kielektroni, X, A na B. Katika spectroscopy ya molekuli, X kwa kawaida huonyesha hali ya kielektroni ya msingi—hali ya nishati ya chini zaidi ya molekuli isiyo na chaji—huku A na B zikiwa hali mbili za kwanza zilizosisimuliwa zinazoonekana katika spectrum. Spectrum ya nishati ya juu katika 6.424 eV haikuonyesha vipengele vingine.

Vipimo vya azimio la juu vilitoa muundo wa kina zaidi wa bendi hizo.

- **2.3429 eV** kwa X state, ambayo ni electron affinity ya neutral CBI;
- **2.5812 eV** kwa A state;
- **3.5246 eV** kwa B state.

Reported experimental kutokuwa na uhakika ni takribani **0.0010 eV** kwa electronic energies na **8 cm⁻¹** kwa vibrational frequencies.

Measured vibrational frequencies zilikuwa karibu lakini si identical:

- X: **681 cm⁻¹**;
- A: **606 cm⁻¹**;
- B: **633 cm⁻¹**.

Namba hizo ni muhimu kwa sababu zinasema kitu kuhusu bonding katika kila neutral state. Bond fupi na strong kwa kawaida hutoa higher stretching frequency; weaker au longer bond huielekea kuishusha. Chemistry haitoi rule hiyo bila caveats kila mara, lakini ni first handle muhimu.

Picha ya zamani inaanza kupata shida wapi

Katika nonrelativistic picha, CBI- ingeonekana kama cousin nzito ya CN-. Ungetarajia filled sigma orbital na filled pi orbitals mbili. Kuondoa electron moja kungeleta neutral states zinazoweza kuasignifiwa katika ordinary sigma/pi lugha.

vipimo hazifuati picha hiyo kwa usafi.

Angular distributions ndiyo tatizo la kwanza. Detector hapa haipimi energy ya electron pekee; pia directions ambako electrons zinatoka. Directional pattern hiyo inafupishwa kwa anisotropy parameter beta. X band ina beta **1.86**, ambayo waandishi wanatafsiri kama dominant p-wave detachment kutoka sigma-type orbital. A na B bands zina beta **-0.73** na **-0.67**, consistent na more pi-like detachment.

Mpaka hapo inaonekana manageable: X ni sigma-like, A na B ni pi-like.

Lakini vibrational structure inakataa assignment ileile. Electron ikiondolewa, molecule inaweza kubaki vibrating; pattern ya vibration peaks huitwa Franck-Condon progression. X na B zina similarly short progressions, wakati A ina longer one. Kama A na B zingekuwa tu two spin-orbit components za ordinary pi-hole state moja, zingetarajiwa kufanana zaidi katika bond-length changes. Badala

yake, B inaonekana kama X katika respect moja na kama A katika nyingine.

Hii ndiyo contradiction ambayo ni rahisi kuficha chini ya diagram. Waandishi wanafanya kinyume: wanaitumia kama clue kwamba diagram si object sahihi tena.

Relativistic description

Kwa atomi nzito, spin na mwendo wa obiti haviwezi kutenganishwa kwa urahisi. Kiasi kinachofaa zaidi kuhifadhiwa ni makadirio ya jumla ya momentum ya angular ya elektroni kwenye mhimili wa molekuli. Makala huonyesha kiasi hiki kwa omega.

Hilo linabadilisha basis ya description. Badala ya kutibu triple bond kama sigma moja na pi mbili orbitals kisha kuongeza spin baadaye, waandishi wanaeleza relevant states kama relativistic Kramers pairs. Kramers pair ni pair ya degenerate spinors inayohitajika na time-reversal symmetry katika mifumo zenye odd number of electrons. Neno ni technical, lakini point ni rahisi: katika relativistic case, natural one-electron objects ni spinors, si ordinary spin-free orbitals ambazo spin inawekwa juu baadaye.

Fully relativistic calculation inatoa pure pi-like $|\omega| = 3/2$ Kramers pair moja na $|\omega| = 1/2$ Kramers pairs mbili zenye substantial sigma/pi mixing. Kwa lugha rahisi, pair moja bado inatenda mostly kama pi component, wakati nyingine mbili haziwezi kubaki cleanly sigma au pi. Hiyo ndiyo “collapse” katika title ya makala: si bonding kutoweka, bali classical sigma/pi separation kuacha kuwa lugha sahihi kwa molecule hii.

Computations si decorative afterthought. Waandishi wanatumia four-component Dirac-Coulomb coupled-cluster mbinu, pamoja na DC-CCSD(T) kwa ground na low-lying states na EOM-IP-CCSD kwa B state. Calculated C-Bi bond length kwa C-Bi- ni **2.022 angstroms**, na computed anion stretching frequency ni **695 cm⁻¹**, karibu na experimental scale. Calculated adiabatic detachment energies zinakubaliana vizuri na measured X na A states na zinaunga mkono relativistic assignment.

B state bado ndiyo delicate one kwa calculation. Calculated vibrational frequency yake hailingani vizuri sawa na experiment, na waandishi wanaiona kama sign kwamba frequency ni sensitive sana kwa degree ya mixing kati ya X na B states. Strong $|\omega| = 1/2$ mixing hiyohiyo ndiyo inayofanya B kuwa na frequency noticeably higher kuliko A. Clean makala haipaswi kuwa cleaner kuliko makala inayoieleza.

Kile ambacho hii haithibitishi

- **Haimaanishi** ordinary sigma na pi bonding hazina matumizi.
- **Haimaanishi** carbon-nitrogen triple bonds, alkynes au light-element textbook examples nyingi zinahitaji kuchorwa upya.
- **Haithibitishi** kila heavy-element bond inatenda kama C-Bi-.
- **Haisemi** C-Bi bond si multiple bond.

- **Haifanyi** relativistic quantum chemistry kuwa optional flourish; katika case hii inahitajika kwa assignment sahihi.
- **Haitengenezi** material mpya au chemical technology mpya yenyewe.

Boundary ndiyo point. Classical bonding lugha hufanya kazi vizuri dhana zake zikiwa approximately true. CBI- ni muhimu kwa sababu dhana hizo zinasukumwa kwa nguvu ya kutosha failure ione kane.

Ushahidi una nguvu kiasi gani?

ushahidi ni strong kwa assignment waandishi wanayofanya.

Experimentally, makala inaunganisha high-resolution cryogenic spectra, vibrational structure na photoelectron angular distributions. Hizo ni complementary constraints: energy spacings pekee zingekuwa weaker; angular distributions pekee weaker; tension kati yake ndiyo inaelekeza relativistic interpretation.

Computationally, waandishi wanatumia fully relativistic four-component mbinu badala ya kuongeza spin-orbit coupling kama small correction baada ya nonrelativistic calculation. Hilo ni muhimu kwa sababu dai hasa ni kwamba spin-orbit coupling si ndogo katika molecule hii.

Match si perfect. B-state vibrational frequency ndiyo notable loose end. makala pia inasoma molecular ion moja, si chemical universe nzima. Lakini kama kipimo linganishi case ya relativistic heavy-element bonding, CBI- ni clean isivyo kawaida: ndogo, experimentally resolved na theoretically tractable.

Kwa nini ni muhimu

Chemistry mara nyingi hufundisha bonding kwa picha. Hilo si weakness. picha nzuri inacompress quantum mechanics kuwa kitu ambacho binadamu anaweza kutumia.

Lakini kila picha ina jurisdiction yake. Sigma/pi triple-bond picha ni ya regime ambapo spin-orbit coupling inaweza kutibiwa kama secondary. Heavy elements zinaweza kutoka regime hiyo. CBI- inaonyesha departure hiyo katika molecule ndogo kiasi kwamba failure inaweza kupimwa na kukokotolewa kwa detail.

Hilo ni muhimu kwa heavy-element chemistry kwa sababu bismuth na neighbours zake si exotic curiosities katika quantum mechanics. Ni maeneo ambapo relativistic athari ni sehemu ya ordinary accounting. Chemists wakitaka kubuni, kuintepret au kupredict bonding karibu na heavy atoms, wanahitaji lugha inayohifadhi quantities sahihi.

Thamani ya makala si kuifanya old picha ione kane foolish. Inafanya kitu muhimu zaidi: inaonyesha hasa wapi old picha inaacha kubeba mzigo.

Muhtasari safi

Kahraman, Hui, Zhang, Ellis, Choi, Peterson na Wang walipima CBi- molecular ion kwa high-resolution cryogenic photoelectron spectroscopy na photoelectron upigaji picha, kisha wakalinganisha spectra na fully relativistic Dirac-Coulomb coupled-cluster calculations. Walipata neutral CBi states tatu zenye adiabatic detachment energies 2.3429, 2.5812 na 3.5246 eV. Spectra na angular distributions haziingii vizuri katika simple nonrelativistic sigma-plus-two-pi triple-bond picha. Badala yake, strong spin-orbit coupling inareorganize bonding kuwa relativistic Kramers pairs: pi-like $|\omega| = 3/2$ pair moja na $|\omega| = 1/2$ pairs mbili zenye sigma/pi mixing. Hii ni moja kwa moja ushahidi kwamba katika very heavy-element triple-bond mfumo, textbook orbital labels zinaacha kuwa conserved lugha bora. Si rejection ya ordinary chemical bonding; ni map sahihi ya sehemu moja ambapo relativity inachukua bookkeeping.

Vyanzo

Kahraman, Hui, Zhang, Ellis, Choi, Peterson and Wang, Relativistic collapse of the classical triple bond in the CBi- molecular ion, *Science* 393, 184-187 (2026)

<https://doi.org/10.1126/science.aei1285>

Kahraman et al., Tabular Data and Computational Details for the CBi Spectroscopic Analysis, Zenodo (2026)

<https://doi.org/10.5281/zenodo.20249312>