

data_bcx1dc_abs

_audit_creation_method SHELXL-97
_chemical_name_systematic
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?
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_chemical_name_common ?
_chemical_melting_point ?
_chemical_formula_moiety ?
_chemical_formula_sum
'H B Ca O5 Si'
_chemical_formula_weight 159.99

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_atom_type_description
_atom_type_scat_dispersion_real
_atom_type_scat_dispersion_imag
_atom_type_scat_source
'H' 'H' 0.0000 0.0000
'International Tables Vol C Tables 4.2.6.8 and 6.1.1.4'
'B' 'B' 0.0013 0.0007
'International Tables Vol C Tables 4.2.6.8 and 6.1.1.4'
'O' 'O' 0.0106 0.0060
'International Tables Vol C Tables 4.2.6.8 and 6.1.1.4'
'Si' 'Si' 0.0817 0.0704
'International Tables Vol C Tables 4.2.6.8 and 6.1.1.4'
'Ca' 'Ca' 0.2262 0.3064
'International Tables Vol C Tables 4.2.6.8 and 6.1.1.4'

_symmetry_cell_setting ?
_symmetry_space_group_name_H-M ?

loop_
_symmetry_equiv_pos_as_xyz
'x, y, z'
'-x, y+1/2, -z+1/2'
'-x, -y, -z'
'x, -y-1/2, z-1/2'

_cell_length_a 4.8332(2)
_cell_length_b 7.6082(2)
_cell_length_c 9.6342(2)
_cell_angle_alpha 90.00
_cell_angle_beta 90.147(2)
_cell_angle_gamma 90.00
_cell_volume 354.267(19)
_cell_formula_units_Z 4
_cell_measurement_temperature 293(2)
_cell_measurement_reflns_used ?
_cell_measurement_theta_min ?

_cell_measurement_theta_max ?
_exptl_crystal_description ?
_exptl_crystal_colour ?
_exptl_crystal_size_max ?
_exptl_crystal_size_mid ?
_exptl_crystal_size_min ?
_exptl_crystal_density_meas ?
_exptl_crystal_density_diffn 3.000
_exptl_crystal_density_method 'not measured'
_exptl_crystal_F_000 320
_exptl_absorpt_coefficient_mu 1.998
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_exptl_absorpt_correction_T_min ?
_exptl_absorpt_correction_T_max ?
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_exptl_special_details

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?
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_diffrn_ambient_temperature 293(2)
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_diffrn_radiation_monochromator graphite
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_diffrn_standards_number ?
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_diffrn_reflns_limit_h_max 8
_diffrn_reflns_limit_k_min -13
_diffrn_reflns_limit_k_max 13
_diffrn_reflns_limit_l_min -16
_diffrn_reflns_limit_l_max 17
_diffrn_reflns_theta_min 3.41
_diffrn_reflns_theta_max 40.12
_reflns_number_total 2192
_reflns_number_gt 1984
_reflns_threshold_expression >2sigma(I)

_computing_data_collection ?
_computing_cell_refinement ?
_computing_data_reduction ?
_computing_structure_solution ?
_computing_structure_refinement 'SHELXL-97 (Sheldrick, 1997)'

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_computing_molecular_graphics    ?
_computing_publication_material ?  

  

_refine_special_details  

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Refinement of F^2^ against ALL reflections. The weighted R-factor wR and  

goodness of fit S are based on F^2^, conventional R-factors R are based  

on F, with F set to zero for negative F^2^. The threshold expression of  

F^2^ > 2sigma(F^2^) is used only for calculating R-factors(gt) etc. and is  

not relevant to the choice of reflections for refinement. R-factors based  

on F^2^ are statistically about twice as large as those based on F, and R-  

factors based on ALL data will be even larger.  

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_refine_ls_structure_factor_coef  Fsqd  

_refine_ls_matrix_type          full  

_refine_ls_weighting_scheme    calc  

_refine_ls_weighting_details  

'calc w=1/[s^2^(Fo^2^)+(0.0200P)^2^+0.0000P] where P=(Fo^2^+2Fc^2^)/3'  

_atom_sites_solution_primary   direct  

_atom_sites_solution_secondary difmap  

_atom_sites_solution_hydrogens geom  

_refine_ls_hydrogen_treatment mixed  

_refine_ls_extinction_method  SHELXL  

_refine_ls_extinction_coef    0.002(5)  

_refine_ls_extinction_expression  

'Fc^*^=kFc[1+0.001xFc^2^|I^3^/sin(2|q)]^-1/4^'  

_refine_ls_number_reflns      2192  

_refine_ls_number_parameters  78  

_refine_ls_number_restraints  0  

_refine_ls_R_factor_all       0.0407  

_refine_ls_R_factor_gt        0.0375  

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_refine_ls_wR_factor_gt      0.0835  

_refine_ls_goodness_of_fit_ref 1.813  

_refine_ls_restrained_S_all  1.813  

_refine_ls_shift/su_max       0.001  

_refine_ls_shift/su_mean      0.000  

  

loop_
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_atom_site_type_symbol
_atom_site_fract_x
_atom_site_fract_y
_atom_site_fract_z
_atom_site_U_iso_or_equiv
_atom_site_adp_type
_atom_site_occupancy
_atom_site_symmetry_multiplicity
_atom_site_calc_flag
_atom_site_refinement_flags
_atom_site_disorder_assembly
_atom_site_disorder_group
Ca Ca 0.99165(4) 0.10619(3) 0.33611(2) 0.00670(7) Uani 1 1 d . .

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Si Si 0.46894(6) 0.26609(4) 0.08439(3) 0.00445(8) Uani 1 1 d . . .
 B B 0.5685(2) 0.41059(15) 0.34075(12) 0.00556(17) Uani 1 1 d . . .
 O1 O 0.24029(18) 0.40110(11) 0.03750(9) 0.00786(14) Uani 1 1 d . . .
 O2 O 0.67035(17) 0.30087(11) 0.45724(8) 0.00670(13) Uani 1 1 d . . .
 O3 O 0.67535(17) 0.33466(11) 0.21047(8) 0.00662(13) Uani 1 1 d . . .
 O4 O 0.31490(18) 0.08794(11) 0.14605(9) 0.00724(14) Uani 1 1 d . . .
 O5 O 0.25877(18) 0.41345(12) 0.33650(9) 0.00769(14) Uani 1 1 d . . .
 H H 0.216(5) 0.463(3) 0.404(3) 0.028(6) Uiso 1 1 d . . .

loop_

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 _atom_site_aniso_U_22
 _atom_site_aniso_U_33
 _atom_site_aniso_U_23
 _atom_site_aniso_U_13
 _atom_site_aniso_U_12
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 Si 0.00571(13) 0.00498(13) 0.00265(12) -0.00033(8) -0.00067(9) 0.00056(9)
 B 0.0066(4) 0.0065(4) 0.0035(4) -0.0003(3) -0.0003(3) -0.0006(3)
 O1 0.0082(3) 0.0087(3) 0.0066(3) -0.0004(2) -0.0020(2) 0.0029(2)
 O2 0.0075(3) 0.0086(3) 0.0040(3) 0.0019(2) 0.0007(2) 0.0012(2)
 O3 0.0073(3) 0.0096(3) 0.0029(3) -0.0017(2) -0.0007(2) 0.0000(2)
 O4 0.0082(3) 0.0055(3) 0.0080(3) 0.0008(2) 0.0016(2) 0.0004(2)
 O5 0.0063(3) 0.0106(3) 0.0062(3) -0.0008(2) -0.0003(2) 0.0011(2)

_geom_special_details

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All esds (except the esd in the dihedral angle between two l.s. planes) are estimated using the full covariance matrix. The cell esds are taken into account individually in the estimation of esds in distances, angles and torsion angles; correlations between esds in cell parameters are only used when they are defined by crystal symmetry. An approximate (isotropic) treatment of cell esds is used for estimating esds involving l.s. planes.

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loop_

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 _geom_bond_atom_site_label_2
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 _geom_bond_site_symmetry_2
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 Ca O1 2.2762(9) 2_645 ?
 Ca O1 2.2803(8) 4_666 ?
 Ca O4 2.4141(9) 1_655 ?
 Ca O2 2.4447(8) . ?
 Ca O5 2.5240(9) 2_645 ?
 Ca O3 2.6104(9) . ?
 Ca O3 2.6577(9) 2_745 ?
 Ca O5 2.6705(9) 1_655 ?
 Si O1 1.5741(9) . ?
 Si O2 1.6474(9) 4_565 ?
 Si O3 1.6543(8) . ?
 Si O4 1.6573(9) . ?

B O4 1.4676(15) 2_655 ?

B O3 1.4761(14) . ?

B O2 1.4818(14) . ?

B O5 1.4976(15) . ?

O1 Ca 2.2762(9) 2_655 ?

O1 Ca 2.2803(8) 4_465 ?

O2 Si 1.6474(9) 4_566 ?

O3 Ca 2.6577(9) 2_755 ?

O4 B 1.4676(15) 2_645 ?

O4 Ca 2.4141(9) 1_455 ?

O5 Ca 2.5240(9) 2_655 ?

O5 Ca 2.6705(9) 1_455 ?

loop_

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_geom_angle_atom_site_label_2

_geom_angle_atom_site_label_3

_geom_angle

_geom_angle_site_symmetry_1

_geom_angle_site_symmetry_3

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O1 Ca O1 77.72(3) 2_645 4_666 ?

O1 Ca O4 133.39(3) 2_645 1_655 ?

O1 Ca O4 107.66(3) 4_666 1_655 ?

O1 Ca O2 81.12(3) 2_645 . ?

O1 Ca O2 86.71(3) 4_666 . ?

O4 Ca O2 144.10(3) 1_655 . ?

O1 Ca O5 73.63(3) 2_645 2_645 ?

O1 Ca O5 142.87(3) 4_666 2_645 ?

O4 Ca O5 77.12(3) 1_655 2_645 ?

O2 Ca O5 111.25(3) . 2_645 ?

O1 Ca O3 114.57(3) 2_645 . ?

O1 Ca O3 135.85(3) 4_666 . ?

O4 Ca O3 93.78(3) 1_655 . ?

O2 Ca O3 56.35(3) . . ?

O5 Ca O3 78.59(3) 2_645 . ?

O1 Ca O3 81.77(3) 2_645 2_745 ?

O1 Ca O3 78.86(3) 4_666 2_745 ?

O4 Ca O3 55.51(3) 1_655 2_745 ?

O2 Ca O3 159.60(3) . 2_745 ?

O5 Ca O3 74.14(3) 2_645 2_745 ?

O3 Ca O3 142.512(13) . 2_745 ?

O1 Ca O5 146.94(3) 2_645 1_655 ?

O1 Ca O5 76.49(3) 4_666 1_655 ?

O4 Ca O5 74.79(3) 1_655 1_655 ?

O2 Ca O5 77.09(3) . 1_655 ?

O5 Ca O5 137.775(19) 2_645 1_655 ?

O3 Ca O5 72.56(3) . 1_655 ?

O3 Ca O5 112.81(3) 2_745 1_655 ?

O1 Si O2 113.84(4) . 4_565 ?

O1 Si O3 115.25(5) . . ?

O2 Si O3 106.69(4) 4_565 . ?

O1 Si O4 108.72(5) . . ?

O2 Si O4 106.29(4) 4_565 . ?

O3 Si O4 105.39(4) . . ?
O4 B O3 107.36(9) 2_655 . ?
O4 B O2 109.00(8) 2_655 . ?
O3 B O2 107.89(9) . . ?
O4 B O5 111.87(9) 2_655 . ?
O3 B O5 109.53(8) . . ?
O2 B O5 111.04(9) . . ?
Si O1 Ca 129.80(5) . 2_655 ?
Si O1 Ca 126.61(5) . 4_465 ?
Ca O1 Ca 102.28(3) 2_655 4_465 ?
B O2 Si 122.79(7) . 4_566 ?
B O2 Ca 100.95(6) . . ?
Si O2 Ca 123.02(4) 4_566 . ?
B O3 Si 122.42(7) . . ?
B O3 Ca 94.13(6) . . ?
Si O3 Ca 118.80(4) . . ?
B O3 Ca 93.02(6) . 2_755 ?
Si O3 Ca 119.04(4) . 2_755 ?
Ca O3 Ca 103.99(3) . 2_755 ?
B O4 Si 127.61(7) 2_645 . ?
B O4 Ca 103.67(6) 2_645 1_455 ?
Si O4 Ca 121.13(4) . 1_455 ?
B O5 Ca 120.27(6) . 2_655 ?
B O5 Ca 118.07(6) . 1_455 ?
Ca O5 Ca 106.06(3) 2_655 1_455 ?

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_refine_diff_density_rms 0.194