

TABLE II. General Cauchy relations for first-, second-, and third-order elastic constants. Also listed are coefficients used in the general expressions for the elastic constants.

Types <i>ijklmn</i>	Elastic constants with Cauchy relations $C_{ij\dots}$	$X_{ij\dots}$	$Y_{ijkl\dots}$	Z_{ijklmn}
First order				
<i>II</i>	C_1, C_2, C_3	1		
<i>JK</i>	C_4, C_5, C_6	0		
Second order				
<i>III</i>	C_{11}, C_{22}, C_{33}	3	$6g_I^2$	
<i>IJJ</i>	$C_{12}=C_{66}, C_{13}=C_{55}, C_{23}=C_{44}$	1	$g_I^2+g_J^2$	
<i>JJK</i>	$C_{15}, C_{16}, C_{24}, C_{26}, C_{34}, C_{35}$	0	$3g_Jg_K$	
<i>IJK</i>	$C_{14}=C_{56}, C_{25}=C_{46}, C_{36}=C_{45}$	0	g_Jg_K	
Third order				
<i>IIII</i>	$C_{111}, C_{222}, C_{333}$	15	$45g_I^2$	$15g_I^4$
<i>IIIIJ</i>	$C_{112}=C_{166}, C_{113}=C_{155}, C_{122}=C_{266}$ $C_{133}=C_{355}, C_{223}=C_{244}, C_{233}=C_{344}$	3	$6g_I^2+3g_J^2$	$g_I^4+6g_I^2g_J^2$
<i>IIJJK</i>	$C_{123}=C_{144}=C_{255}=C_{366}=C_{456}$	1	$g_I^2+g_J^2+g_K^2$	$g_I^2g_J^2+g_I^2g_K^2+g_K^2g_I^2$
<i>JJJJK</i>	$C_{115}, C_{116}, C_{224}, C_{226}, C_{334}, C_{335}$	0	$15g_Jg_K$	$10g_J^3g_K$
<i>JJJKK</i>	$C_{126}=C_{666}, C_{135}=C_{555}, C_{234}=C_{444}$	0	$9g_Jg_K$	$3g_Jg_K(g_I^2+g_K^2)$
<i>IIIIJK</i>	$C_{114}=C_{156}, C_{225}=C_{246}, C_{336}=C_{345}$	0	$3g_Jg_K$	$6g_I^2g_Jg_K$
<i>IIJJJK</i>	$C_{125}=C_{146}=C_{566}, C_{124}=C_{256}=C_{466}$ $C_{134}=C_{356}=C_{455}, C_{136}=C_{145}=C_{556}$ $C_{235}=C_{346}=C_{445}, C_{236}=C_{245}=C_{446}$	0	$3g_Jg_K$	$g_Jg_K(3g_I^2+g_J^2)$

elastic constants. For convenience, the $X_{ij\dots}$, $Y_{ijkl\dots}$, and Z_{ijklmn} coefficients, which are used in Eqs. (17)–(19) to determine $C_{ij\dots}^{99}$, have also been listed.

Calculations have been completed for seven cubic structures, namely simple cubic, fcc, bcc, NaCl type, CsCl type, diamond, and zinc blende. The first-, second-, and third-order elastic constants with Cauchy relations for these structures are

$$C_1; C_{11}, C_{12}=C_{44};$$

and

$$C_{111}, C_{112}=C_{155}, C_{123}=C_{144}=C_{456}.$$

The electrostatic contributions to these constants

are listed in Table III in units of $Z^2 e^2/a^4$, where a is the lattice parameter.

Four hexagonal structures, namely simple hexagonal, hcp, WC type, and wurtzite, have also been considered. For these structures, the elastic constants with Cauchy relations are

$$C_1, C_3; C_{11}, C_{33}, C_{12}=C_{66}=\frac{1}{3}C_{11}, C_{13}=C_{44};$$

and

$$C_{111}, C_{222}, C_{112}=C_{166}=\frac{1}{5}(-2C_{111}+3C_{222}),$$

$$C_{122}=C_{266}=\frac{1}{5}(3C_{111}-2C_{222}),$$

$$C_{333}, C_{113}=C_{155},$$

TABLE III. Electrostatic Brugger elastic constants for cubic structures. Entries are in units of $Z^2 e^2/a^4$, where a is the lattice constant.

	simple cubic	fcc	bcc	NaCl type	CsCl type	diamond	zinc blende
<i>U</i>	-1.418 648 740	-9.169 724 148	-3.639 233 450	-13.980 516 757	-2.035 361 509	-21.547 192 177	-15.131 704 416
C_1	0.472 882 913	3.056 574 716	1.213 077 817	4.660 172 252	0.678 453 836	7.182 397 392	5.043 901 472
C_{11}	-0.143 189 083	-6.849 873 626	-2.697 885 714	-25.108 469 175	2.125 129 382	-23.345 777 443	-4.053 717 061
C_{12}	-0.637 729 828	-1.159 925 261	-0.470 673 868	5.563 976 209	-2.080 245 446	0.899 292 633	-5.538 993 678
C_{111}	-1.747 954 699	34.065 579 503	9.235 126 065	164.229 593 204	-16.226 944 863	119.055 300 863	17.207 017 149
C_{112}	1.231 950 057	0.091 894 314	2.127 151 252	-19.343 623 664	2.800 648 978	-1.163 206 824	1.530 784 079
C_{123}	0.724 749 027	5.615 837 678	-1.900 933 165	10.867 366 283	4.799 929 272	-2.170 049 518	24.633 400 230

TABLE IV. Electrostatic Brugger elastic constants for hexagonal structures with ideal structure parameters. Entries are in units of $Z^2 e^2/a^4$, where a is the lattice constant.

	simple hexagonal ($c/a=1$)	hcp ($c/a=\sqrt{\frac{8}{3}}$)	WC type ($c/a=\sqrt{\frac{8}{3}}$)	wurtzite ($c/a=\sqrt{\frac{8}{3}}$, $u=\frac{3}{8}$)
U	-1.729 575 102	-2.292 340 210	-0.873 690 983	-3.790 469 934
C_1	0.533 603 351	0.764 635 593	-0.577 163 952	1.237 266 468
C_3	0.662 368 399	0.763 069 024	2.028 018 888	1.315 936 998
C_{11}	-0.605 973 676	-1.288 353 922	2.966 252 428	-2.637 875 499
C_{33}	-0.401 414 892	-1.137 003 974	-1.637 033 899	-3.558 546 850
C_{12}	-0.201 991 225	-0.429 451 308	0.988 750 809	-0.879 291 833
C_{13}	-0.792 845 153	-0.576 101 549	-2.223 511 382	-0.194 632 072
C_{111}	2.324 128 642	5.026 701 204	-15.767 553 741	12.580 565 280
C_{222}	-0.532 222 756	2.217 541 686	-19.362 925 160	2.779 206 544
C_{333}	-0.708 777 482	5.511 268 055	2.608 789 547	26.442 451 932
C_{112}	-1.248 985 111	-0.680 155 470	-5.310 733 599	-3.364 702 185
C_{122}	1.607 366 288	2.129 004 048	-1.715 362 181	6.436 656 550
C_{113}	1.954 724 848	2.095 223 879	6.247 025 199	3.973 514 399
C_{123}	0.651 574 949	0.698 407 960	2.082 341 733	1.324 504 800
C_{133}	1.357 925 970	0.086 875 908	2.788 189 975	-4.324 858 840

$$C_{123} = C_{144} = C_{366} = C_{456} = \frac{1}{3} C_{113}, \quad C_{133} = C_{344}.$$

Electrostatic contributions to these elastic constants for ideal structure parameters are listed in Table IV in units of $Z^2 e^2/a^4$. For all the hexagonal structures, our choice of Cartesian axes (in Miller-Bravais indices) has been 1 axis = $[10\bar{1}0]$; 2 axis = $[\bar{1}2\bar{1}0]$; and 3 axis = $[0001]$. It should be mentioned that, for the hcp structure, Cousins⁵ has chosen basal axes rotated 90° with respect to ours. Thus, his C_{111} corresponds to our C_{222} , etc. However, our choice of axes is consistent with the only two sets of measured third-order elastic constants for hcp metals.^{14,15} Also the different choice of axes eliminates the discrepancy, noted by Naimon *et al.*,¹⁶ in certain Fuchs's constants calculated by Cousins.⁴

The results presented in Tables III and IV represent a higher degree of accuracy than those of earlier calculations. However, our results are essentially in agreement with those reported earlier (fcc,^{1-3,9} bcc,^{1-3,6,9} NaCl and CsCl,⁷ zinc blende,⁸ and hcp⁵). Also, all results were checked independently by doing a Fuchs-type calculation for each structure [Eq. (11) with $\lambda = (\Omega_0')^{1/3}$]. Other useful checks are the relations

$$C_{ii}^{es} = -U_{es}, \quad C_{ijij}^{es} = 3U_{es}, \quad C_{ijjkh}^{es} = -15U_{es},$$

where, as usual, repeated indices are to be summed. These can be easily derived, for example, by relating Fuchs and Brugger constants. All calculations were performed on IBM 360 and Xerox Sigma 5 computers. Convergence of all sums was such that the maximum error in the tabulated constants was $\pm 1 \times 10^{-12}$. The subroutine

for the complementary error function, necessary to generate the $\Phi_{-1/2}$ functions, was from an IBM routine with a relative error of less than 4×10^{-16} .

IV. GENERAL EXPRESSIONS FOR INTERNAL-STRAIN DERIVATIVES

In order to obtain the internal-strain contribution to the elastic constants, it is sufficient to know the energy density of the homogeneously deformed lattice as a function of both external and internal strains (see Appendix A). However, since the internal strains are determined from the total energy density, the electrostatic internal-strain contribution to the elastic constants cannot be obtained directly. It is still possible, though, to determine electrostatic internal-strain derivatives of the form

$$U_{ijkl\cdots}^{(\nu\mu)\cdots} = \left(\frac{\partial^n U_{es}'}{\partial \bar{w}_\nu(\nu) \partial \bar{w}_\mu(\mu) \cdots \partial \eta_{ij} \partial \eta_{kl} \cdots} \right)_{\bar{\eta}=0, \bar{w}=0} \quad (26)$$

where $\bar{w}(\nu)$ is the internal strain associated with the ν th ion of the unit cell ($\nu=0, 1, \dots, s-1$). These derivatives would be combined with those arising from other terms of the energy density, thus resulting in the total internal-strain contribution to the elastic constants.

Using the method of homogeneous deformation, derivatives of $U_{es}'(\bar{\eta}, \bar{w}(\nu))$ can be easily performed (see Appendix B). Introducing the dimensionless parameter $\bar{t} = [\bar{t}(\nu) - \bar{t}(\mu)] / \Omega_0^{1/3}$, as well as $\bar{r} = \bar{R}(\nu_0) / \Omega_0^{1/3}$ and $\bar{g} = \Omega_0^{1/3} \bar{G}(h) / 2\pi$, the expressions for the electrostatic internal-strain derivatives through third-order are