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# A New Approach to Quantum-Statistical Mechanics

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A new method of calculating the grand partition function of many-body system is developed, adopting extensively the techniques of calculus in quantum field theory. It is shown that the grand partition function, which is a trace of the density matrix expressed in terms of field operators, can be evaluated in a way almost parallel with the evaluation of the vacuum expectation value of the S.matrix in quantum field theory, provided that appropriate modifications in notation and definitions are made. As an example, the method is applied to electron-phonon system. Further, basing on this new formalism, various non-perturbational methods are discussed.

#### §1. Introduction

Stimulated by the studies of cooperative phenomena in quantum statistical system such as ferro and antiferromagnetism, superconductivity, the A-transition in liquid helium etc., various methods for the calculation of the partition function of many-body system with interaction have been put forward by many authors. These methods of attack have each its own merit, and in some cases they have been fairly successfully applied to practical problems. For instance, Kubo established an expansion theorem of the density matrix and applied it to ferro and antiferromagnetism.1> Schafroth, in his theory of Meissner effect in superconductors, derived a formula in which the density matrix was expressed in powers of the interaction Hamiltonian.2> Essentially the same formula for the expansion of the density matrix was also obtained by a different method by Chester, who made use of it to discuss the Bose-Einstein condensation of imperfect Bose gas. > A quite different way than others to handle the density matrix was invented by Feynman (the method of integral over trajectories), and he applied it to the problem of liquid helium<sup>4</sup>> More recently, Friedman and Butler introduced another technique of manipulating the density matrix and thereby discussed the transition in liquid helium quantitatively.5>

Generally speaking, however, it seems that major efforts have been made so far to overcome the difficulties encountered in treating the interaction in many-body system, so that there remain unsolved difficulties in taking account of the effect of statistics, especially for Fermion system such as electrons in a superconductor and liquid helium 3, in both of which the role of Fermi statistics seeems to be important. To remedy this point, it may be promising to use the number representation of the second quantization theory for the calculation of the trace of the density matrix. From this view point Ichimura developed a method of expanding the grand partition function in powers of the coupling constant using the number representation.<sup>6</sup> It appears, however, to the present author that his method is unsatisfactory

in the following two points : First it will not be practical in evaluating higher order corrections, because troublesome calculation of an enormous number of terms are needed; Therefore, application of this method will be restricted only to cases in which the effect of higher order perturbations is unimportant. Secondly, while the number representation may be most convenient to take into account the effect of statistics, it has such a defect that it is difficult to treat by this method the problem in confi<sub>g u</sub> ration space, namely, it is not easy by this method to utilize physical pictures connected with the confi<sub>g u</sub> ration space. For instance, one cannot utilize the quantity such as the molecular distribution function, which has been useful for the understanding of the cooperative phenomena in classical system.

In this paper we shall present a new approach which seems to be free from the above mentioned shortcomings of the n-representation. We introduce explicitly the quantized field of particles and utilize the various techniques of operator calculus in quantum field theory as far as possible in evaluating the quantum-statistical average of the field quantities. In § 2 and § 3 we give a general formulation of our theory for an example of electron-phonon system. Various results obtained by means of this new method for electron-phonon system are illustrated in § 4. In § 5 non-perturbational treatments are discussed, starting from the formulation given in § 2 and § 3. The last section is devoted to a possible extension of our method to other systems.

### § 2. General formulation

We suppose that the Hamiltonian of a system in question can be divided into two parts

$$H = H0 + H_{1}$$
(2.1)

each of which is expressed in terms of fi.eld quantity as

$$H_{0}^{-} = \int H (\mathbf{x}) d^{3}\mathbf{x},$$

$$H_{1}^{-} i H_{1}(\mathbf{x}) d^{8}\mathbf{x}.$$
(2.2)

We shall call Ho(x) the Hamiltonian density of free field and  $H_1(x)$  the density of interaction Hamiltonian. The density matrix of a canonical ensemble  $p = \exp(-\{IH\})$  bas to satisfy the Bloch equation

$$-\partial \rho / \partial \beta = (H_0 + H_1) \cdot \rho, \quad \{l = l/kT.$$
(2.3)

If we put

$$\exp(- \{IH\} = \exp(- \{1H0\} \cdot S(\{I\}), (2.4)\}$$

the equation for S(n) becomes

$$-\partial S(\beta)/\partial \beta = H_1(\beta) \cdot S(\beta), \qquad (2.5)$$

where

$$H_1(t) = \exp(tH_0) \cdot H_1 \cdot \exp(-tH_0). \qquad (2 \cdot 6)$$

The solution of (2.5) with initial condition S(O) = 1 may be written as

$$S(\beta) = \sum_{n=0}^{\infty} (-1)^n \int_0^{\beta} dt_1 \int_0^{t_1} dt_2 \int_0^{t_2} \cdots \int_0^{t_{n-1}} dt_n H_1(t_1) H_1(t_2) \cdots H_1(t_n)$$
  
= 
$$\sum_{n=0}^{\infty} (-1)^n / n! \int \cdots \int P[H_1(t_1) \cdots H_1(t_n)] dt_1 \cdots dt_n, \qquad (2 \cdot 7)$$

where P is an ordering operator<sup>7</sup> which re-analogs the operators in the bracket in such an order that the arguments t in them are decreasing in magnitude, that is,

$$P[H_1(t_1)\cdots H_1(t_n)] = H_1(t_1')H_1(t_2')\cdots H_1(t_n')$$
  
$$t_1' > t_2' > \cdots > t_n'.$$
(2.8)

If we regard

$$\exp(tH_0) \cdot H_1(\mathbf{x}) \cdot \exp(-tH_0) = H_1(\mathbf{x}, t) \equiv H_1(\mathbf{x})$$
(2.9)

as an operator in the four dimensional space with coordinates x = (x, t), then (2 •7) can be put into another form

$$\mathcal{S}(\beta) = \sum_{n=0}^{\infty} (-1)^n / n! \int \cdots \int P[H_1(x_1) \cdots H_1(x_n)] d^4 x_1 \cdots d^4 x_n, \qquad (2 \cdot 10)$$

in which  $d^4x = d^3x dt$  and the integrations are to be taken over the whole volume with respect to  $x_1$  and over the range (O, /3) with respect to  $t_1$ . P is now an operator arranging the operators in the bracket in such an order that the fourth components of coordinates in them are decreasing in magnitude.

The grand partition function of the system is defined by

$$B = Tm[exp(-aN-/\Im H)], \qquad (2.11)$$

where N is an operator representing the total number of particles, say, of electrons, and a a selector which is related to the chemical potential per one particle  $\mu$  through

$$a = -(\Im \mu. \tag{2.12})$$

Introducing the following notation

(

we rewrite  $(2 \cdot 11)$  in the following forms:

$$B / S_{0^{-}}$$
 (S(/ $\Im$ )) (2 · 14a)

$$= I + f1 + f2 + f3 + \cdot ..$$
 (2.14b)

$$=_{\exp}(C+C+C; +C; \bullet, \bullet), \qquad (2\cdot 14c)$$

where

$$\xi_n = (-1)^n / n! \int \cdots \int \langle P[H_1(x_1) \cdots H_1(x_n)] \rangle d^4 x_1 \cdots d^4 x_n.$$
 (2.15)

The relations between f n's and  $C_{i}$ 's are essentially the same as those between the momen and Thiele's semi-invariants in the theory of probability, that is,

$$f_{\Pi} = \prod_{\substack{2:,lmz = n \\ i}} \Pi(C1) \tau 1/\tau 1 !,$$

$$C_{k} = \sum_{\substack{1 \\ j \\ k}} (-1)^{(\Sigma m_{l}-1)} (\sum_{i} m_{i}-1)! \quad \Pi(\hat{\varsigma}_{i})^{m_{l}}/m_{i}!.$$
(2.16)

3 or represents the grand partition function of the free system in which the interactions are absent. (...) means the quantum-statistical average of a given field quantity denoted by dots referred to the thermal equilibrium realized in the free system.

In order to facilitate the explanation of our further analyses, we consider hereafter as an example the electron-phonon system, whose Hamiltonian is given  $b_v^{-8}$ 

$$H = H 0 + H_{1},$$
 (2.17a)

$$Ho = \mathop{\ddot{e}k}_{k} a: a_{,,+} + \frac{1}{2} \operatorname{Q(I)}_{W} \quad w (\mathbf{b}_{,,+} \mathbf{b}_{,,+} \mathbf{b}_{,+} \mathbf{b}_{,+})$$
(2.17b)

$$H_{1} = g \sum_{k,w} (\hbar \omega_{w}/2V)^{1/2} (a_{k+w}^{*} a_{k} b_{w} + a_{k-w}^{*} a_{k} b_{w}^{*}) + g'/2 f_{::(IJ,V Cb.;:bn, + bmb; + bn,b-w+b.:b!;.,)}, \qquad (2 \cdot 17c)$$

where a f and a<sub>k</sub> represent, respectively, the creation and annihilation operators of the electron with momentum k and energy  $\ddot{\mathbf{e}}_k$ ,  $\mathbf{b}_w$  and  $\mathbf{b}_w$  are the corresponding operators for the phonon with momentum w and energy fur, g and g' are, respectively, a coupling constant and a renormalization constmt given by

$$g = (VC^2/NMs^2)^{1/2}, \quad g' = (s/-s^\circ)/2s^\circ.$$
 (2.18)

The meaning of the symbols appearing in  $(2 \cdot 18)$  is as follows: V is the volume of the system, M the mass of an ion, N the total numbers of ions, C the usual interaction constant betwen electron and lattice,  $s_0$  the sound velocity of free phonons.  $s_0$  is ger..erally different from the real velocity s because there are interactions between electrons and phonons.

Now let us define quantized wave functions of electrons and phonons by

$$\Phi^* (\mathbf{x}) = \frac{\mathbf{v} \cdot 1/2}{k} a_{ik} e^{i\mathbf{k}\cdot\mathbf{x}},$$

$$\psi(\mathbf{x}) = V^{-1/2} \sum_{k} a_{k} e^{i\mathbf{k}\cdot\mathbf{x}},$$

$$\varphi(\mathbf{x}) = \sum_{k} (\hbar w s/2V)^{1/2} (b_{w} e^{iw\cdot\mathbf{x}} + b_{w}^* e^{-iw\cdot\mathbf{x}}).$$
(2.19)

Making use of the commutation relations

. .

$$[a_k, a_{k'}^*]_+ = \delta_{k,k'}, \quad [b_w, b_{w'}^*]_- = \delta_{w,w'},$$

four dimensional fields  $\phi^*(x) = \exp(tH_0)\phi^*(x)\exp(-tH_0)$  etc. are easily shown to become

$$\Phi^*(\mathbf{x}) = \mathbf{v} - 1 \cdot 1 \cdot 2 \quad a: \quad e^{-i\mathbf{k} \cdot \mathbf{x} \cdot \mathbf{H} \mathbf{k} \cdot \mathbf{k}}$$

A New Approach to Quantum-Statistical Mechanics 355

$$\psi(\mathbf{x}) = V^{-1/2} \sum_{k} a_{k} e^{ik \cdot \mathbf{x} - \varepsilon_{k} t}, \qquad (2 \cdot 20)$$

$$\varphi(\mathbf{x}) = \sum_{w} (\hbar w s/2V)^{1/2} (b_{w} e^{iw \cdot \mathbf{x} - \tilde{h} w s t} + b_{w}^{*} e^{-iw \cdot \mathbf{x} + \tilde{h} w s t}).$$

Furtherm.ore one can verify by a direct calculation that

$$e^{tH_0}H_1e^{-tH_0} \equiv H_1(t) = g \int \psi^*(x)\psi(x)\varphi(x)d^3x + g' \int \varphi(x)\varphi(x)d^3x,$$
(2.21)

or

 $H_1(x) = q\phi^*(x)\phi(x)ep(x) + q'ep(x)ep(x).$ 

From  $(2 \cdot 14)$ ,  $(2 \cdot 15)$  and  $(2 \cdot 21)$ , it can be seen that those which we have to know are rules for calculating the averages such as

$$\langle P[\psi^*(\mathbf{x}_1)\psi(\mathbf{x}_1)\psi^*(\mathbf{x}_2)\psi(\mathbf{x}_2)\cdots\psi^*(\mathbf{x}_n)\psi(\mathbf{x}_n)] \rangle$$

$$\langle P[\psi(\mathbf{x}_1)\psi(\mathbf{x}_2)\cdots\psi(\mathbf{x}_n)] \rangle.$$

$$(2.22)$$

and

 $\langle P[\varphi(x_1)\varphi(x_2)\cdots\varphi(x_n)]\rangle$ 

We want to emphasize here that a remarkable similarity exists between the evaluation of 3 / 3 o and that of the vacuum expectation of the so-called S-matrix in quantum field theory. In fact, it will be shown in the next section that all the rules of calcul.:.tions of the vacum expectation of the field quantities in quantum field theory can be used in the present case with only slight modifications.

# §3. Computation rules<sup>7-9>10</sup>>

It will be found convenient in later analyses  $t_0$  use in place of the operator P in (2.22) another operator T defined  $\mathbf{b}_{\mathbf{v}}^{9>}$ 

$$T = \delta_p P, \qquad (3 \cdot 1)$$

where  $\mu_p$  takes 1 or - 1 according as the character of the permutation of the electron operators involved is even or odd in going from the written order to the one re-arranged by P. Of course it holds that

$$P[\mathfrak{H}(\psi^*\psi)] = T[\mathfrak{H}(\psi^*\psi)], \qquad (3.2)$$

if  $S_j(\phi^*\phi)$  is any functional of a product of  $\phi^*(x_i)\phi(x_i)$  is as in (2.22). Now let us find the computation rules for  $(T[Sj(\phi^*\phi)])$  and (PIII(ep)]), where (ep) in any functional of a product of  $\varphi(x)$ 's. In the first place, we decomp:>se  $\phi^*(x)$ ,  $\phi(x)$  and 9(x) into two parts, respectively according to

$$\phi^{*}(x) = \Phi(x) + \phi\phi(x), 
\phi(x) = \phi + (x) + \phi - (x), 
\phi(x) = 'f!_{+}(x) + 'P - (x).$$
(3.3)

For a given product X  $_1$ X  $_2$   $\cdots$  X  $_{,,}$ , where X  $_i$  is any one of the components introduced in (3.3), we define an Nproduct by

$$N[X_{1}X_{2}\cdots X_{n}] = \delta_{p}X_{1'}X_{2'}\cdots X_{n'}, \qquad (3 \ 4)$$

in which the right hand side is a product of the same factors  $X_1X_2 \cdots X\pi$  but ordered in such a manner that all the operators with suffix - stand to the left of all the operators with suffix + and, among the electron operators with the same suffix, all the operators with \* stand to the left of those without  $* \quad iJ$  determines the sign of the permutation in the same way as in  $(3 \cdot 1) \cdot *$  For instance,

$$\begin{split} N[sb^*(x) \ \phi(x')] &= N[\ \Phi t \ (x) \ \Phi_+(x') \ + \Phi Hx)\phi_-(x') \ + \Phi'!. \ (x) \ \Phi_+(x') \ + \Phi'!. \ (x)\phi_-(x')] \\ &= \Phi ! \ (x) \ \Phi_+(x') \ - \phi_-(x')\phi ! \ (x) \ + \Phi'!. \ (x)\Phi_+(x') \ + \Phi'!. \ (x) \ \Phi_-(x')' \\ N[\phi^*(x_1) \ \Phi^*(x_2)\phi(x_3)] &= \phi'!. \ (xi) \ \phi! \ (x_2)\phi_-(x_3) \ + \phi'!. \ (x_1) \ \phi! \ (x_2) \ \Phi_+(x_3) \\ &- \phi'!. \ (x_1) \ \phi_-(x_3) \ \Phi t \ (x_2) \ + \phi! \ (xi) \ \Phi: \ (x_2) \ \phi_+(x_3) \ + \phi! \ (x_2) \ \phi_-(x_3) \ \Phi: \ (x_1) \\ &- \phi'!. \ (x_2) \ \Phi t \ (x_1) \ \Phi_+(x_3) \ + \Phi-(x_3) \ \Phi Hx_1) \ \Phi! \ (x_2) \ + \Phi! \ (x_1) \ \phi t \ (x_2) \ \Phi_+(x_3), \\ N[So \ (x_1) \ So(x_2) \ ] &= So- \ (x_1) \ So-(x_2) \ + \ So-(x_1) \ So-(x_2) \ So+(x_1) \ + \ So+(x_1) \ So+(x_2), \\ N[So \ (x_1) \ So(x_2) \ So(x_3) \ ] &= So-(x_1) \ So-(x_2) \ So+(x_3) \ + So-(x_1) \ So-(x_3) \ So+(x_3) \ + \ So-(x_1) \ So-(x_2) \ So+(x_3) \ + \ So-(x_1) \ So+(x_2) \ So+(x_1) \ So+(x_2) \ So+(x_3), \\ &+ \ So-(x_1) \ So+(x_2) \ So+(x_3) \ + So-(x_3) \ So+(x_1) \ So+(x_2) \ So+(x_1) \ So+(x_2) \ So+(x_3), \\ &+ \ So-(x_2) \ So+(x_1) \ So+(x_3) \ + So-(x_3) \ So+(x_1) \ So+(x_2) \ So+(x_1) \ So+(x_2) \ So+(x_3), \\ &+ \ So-(x_2) \ So+(x_1) \ So+(x_3) \ + \ So-(x_3) \ So+(x_1) \ So+(x_2) \ + \ So+(x_1) \ So+(x_2) \ So+(x_3), \\ &+ \ So-(x_2) \ So+(x_1) \ So+(x_3) \ + \ So-(x_3) \ So+(x_1) \ So+(x_2) \ + \ So+(x_1) \ So+(x_2) \ So+(x_3), \\ \end{array}$$

and so on. Then we can show that an arbitral y T product of electron and phonon field operators is always converted to its corresponding N product through a simple relation. For  $T[\phi^*(x) \phi(x')]$ , it is easy to verify by a direct calculation that

$$T[\phi^*x)\phi(x')] = \frac{N[\phi^*(x)\phi(x')] + [\Phi H x), \ \phi_{-}(x')]_{+}}{1 N[\phi^*(x)\phi(x')] + [\Phi!(x), \ \phi_{-}(x')]_{+} - [\Phi^*(x), \ \phi(x')]_{+} \ t < t'} (3 \cdot 5)$$

For P[So(x)So(x')], we get

$$P[S_{0}(x) S_{0}(x')] = \frac{N[S_{0}(x) S_{0}(x')] + [S_{0+}(x) S_{0-}(x')] - t > t'}{1 N[S_{0}(x) S_{0}(x')] + [S_{0+}(x'), S_{0-}(x)] - t < t'}.$$
(3.6)

Thus if we define two functions  $S(x-x^{\prime})$  and  $D(x-x^{\prime})$  by

$$S(x-x') = \frac{[\Phi!(x), \Phi_{-}(x')]_{+}}{1 \ [\Phi t(x), \phi_{-}(x')]_{+-} \ [\Phi^{+}(x), \phi(x')]_{+}} \quad t > t'}$$
(3.7)

<sup>\*)</sup> Note that there exists a slight difference between our definition of N-product and that wed in quantum field theory. This difference arises from the fact that  $\Phi_* \overset{*}{\ast}(x)$  and  $\Phi_*(x)$  (or  $\Phi_*^*(x)$  and  $\Phi_-(x)$ ) do not necessarily anti-commute with each other, and ther, fore we have to take care of their order in the product.

A New Approach to Quantum-Statistical Mechanics

$$D(x-x') = \begin{cases} \text{So+}(x), \text{ So-}(x') J- t > t' \\ \text{So+}(x'), \text{ So-}(x) J- t < t', \end{cases}$$
(3.8)

the Tproduct with two factors is expressed as a sum of the corresponding N-product and S or D function :

$$T[\phi^*(x) \phi(x')] = N[\phi^*(x) sb(x')] + S(x-x') , \qquad (3.9)$$

$$P[sc>(x)sc>(x) J=N[sc-(x) So(x)] + D(x-x') .$$
(3.9b)

Trying similar calculations for the T-products with more factors than two, we are led to a conclusion that any T-product can be expressible as a sum of terms, each of which is composed of an N-product multiplied by S or D functions. More correctly, this statement will be mathematically expressed in the following two lemmas  $^{:10J}$ 

Lemrna I For any product of SO(x)'s denoted by (SO), it holds that

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$$T[\mathfrak{F}(\varphi)] = N[\mathfrak{F}(\varphi')], \qquad (3.10)$$

where SO(x) is defined by

$$\operatorname{So}(x) = \operatorname{So}(x) + \int d^4 x' D(x - x^{l}) \, \partial/\partial_{\mathfrak{D}}(x') \, . \tag{3.11}$$

 $\partial/\partial_{\mathfrak{D}}(x)$  is an operator characterized by the commutation relation,

$$[\partial/\partial_{\mathfrak{B}}(x), \ \mathrm{SC}(x')J = \partial(x-x'), \qquad (3.12)$$

that is, an operator representing functional differentiation with respect to  $S_{1}(x)$ .

Lemma II For any product of  $\phi^*(x)$ 'S and  $\phi(x)$ 'S, which is denoted by S';  $\phi^*\phi$ , it holds that

$$T[S'; (\phi^* \phi)] = N[S](\phi^* \phi')], \qquad (3.13)$$

where  $\phi^{*'}(x)$  and  $\phi'(x)$  are defined by

$$\phi^{*'}(x) = \phi^{*}(x) + \int d^{4}x' S(x - x') \partial/\partial \phi(x'),$$
  

$$\phi'(x) = \phi(x) - \int d^{4}x' S(x' - x) \partial/\partial \phi^{*}(x'),$$
(3.14)

respectively.  $\partial/\partial \phi^*(x)$  and  $\partial/\partial \phi(x)$  are operators characterized by the following commutation relations :

A proof of these Iemmas will be given in appendix.  $(3 \cdot 9)$  are the special cases of  $(3 \cdot 10)$  or  $(3 \cdot 13) \cdot$  It should be noted that the above results are valid for an arbitrary choice of the manner of decomposition  $(3 \cdot 3)$ . We can, therefore, decompose the field operators in such a way that the resulting computation rules becomes as simple as possible. A possible good choice will be such as to make the averages of N-products  $(N[\phi^*(x)\phi(x')])$  and

(N[So(x)So(x')]) vanish:

$$(N[\phi^*(x)\phi(x')]) = 0, (N[So(x)g(x')]) = 0.$$
 (3.16)

This choice yields for S(x-x') and D(x-x') the results of the form:

$$S(x-x') = \begin{cases} V^{-1} \sum_{k} f_{k} e^{-ik \cdot (x-x') + \varepsilon_{k}(t-t')} & t > t', \\ V^{-1} \sum_{k} (f_{k}-1) e^{-ik \cdot (x-x') + \varepsilon_{k}(t-t')} & t < t', \end{cases}$$

$$D(x-x') = \bigoplus_{u} (hWS/2V) [(N_{w}+1) e \cdot u \cdot (x-x') - hwsJH^{-1}] + N_{w} e^{-iw \cdot (x-x') + hws[t-t']}], \qquad (3.17b)$$

where [k and Nw represent the average numbers of free electrons with momentum k and of free phonons with momentum w in thermal equilibrium at temperature T, namely

$$fk = (a/:ak) = (e^{HEk} + 1) - 1$$
  
Nu. = (b.Jb<sub>w</sub>) = (e<sup>MW</sup> - 1) - 1 (3 · 18)

(3.17) are readily proved with the help of (3.9), (3.16) and (3.20). Although the averages of Nproducts of higher order do not necessarily vanish, it happens that they are such small quantities that their contributions to the grand partition function can be ignored in the limit of N-H $\Pi$ , V-OO (keeping N/V as a constant). To see this fact, we shall consider ( $T[\phi^*(x)\phi(x)\phi^*(x')\phi(x')]$ ) as an example. According to the lemma II, we get

$$(T[\phi^*(x)\phi(x)\phi^*(x')\phi(x')]) = (N[\phi^{*'}(x)\phi'(x)\phi^{*'}(x')\phi'(x')])$$

$$= (N[\phi^*(x)\phi(x)\phi^*(x')\phi(x')]) - S(x-x') (N[\phi^*(x')\phi(x)])$$

$$-S(x'-x) (N[\phi^*(x)\phi(x')]) + S(x-x) (N[\phi^*(x')\phi(x')])$$

$$+S(x'-x') (N[\phi^*(x)\phi(x)]) - S(x-x')S(x'-x) + S(x-x)S(x'-x')$$

$$= (N[\phi^*(x)\phi(x)\phi^*(x')\phi(x')]) - S(x-x')S(x'-x)$$

$$+S(x-x)S(x'-x'), \qquad (3 \cdot 19)$$

in which the condition  $(N[\phi^*(x)\phi(x')])=O$  has been used. On the other hand, if one compute the left-hand side of (3 •19) directly, it will follow that

The average (altaiat.a,.) vanishes for all values of  $(k \ l \ m \ n)$  except for the following three cases :

b) 
$$k=n \ l=m$$
,  
c)  $k=l=m=n$ .

Referring to the definition of S(x-x') given by  $(3\cdot 17a)$ , one can easily see that case a) gives us the term S(x-x)S(x'-x') in the right hand side of  $(3\cdot 10)$ , and the case b) the term S(x-x')S(x'-x), so that

$$(\mathbb{N}[\phi^{*}(\mathbf{x})\phi(\mathbf{x})\phi^{*}(\mathbf{x}')\phi(\mathbf{x}')]) = \mathcal{V}^{-2}(a/a \quad kaf \ a_{k}), \qquad (3 \cdot 21)$$

which is, however, smaller than the other two terms by a factor 1/V, owing to the contraction of the summation over momenta from double to single. The same reasoning prevails for all the averages of Nproducts, and the average of an Nproduct with 2n factors is generally shown to be a quantity of the order of  $(1/v^n - 1)$  if the c-number terms composed of S(x-x') and D(x-x') alone are regarded as the quantities of the order of unity. Thus, on disregarding all the averages of Nproducts, we are led to a simple computation rule for  $(T | H_1(x_1) \cdots H_1(x_m)]$ , that is;

(1) substitute for every field operator  $\phi^*(x)$ ,  $\phi(x)$  and So(x) the quantities  $\phi^{*'}(x)$ ,  $\phi'(x)$  and  $\eta'(x)$  defined by (3.14) and (3.15) respectively,

(2) perform the operations indicated by  $\alpha/\alpha \phi^*(x)$ ,  $\alpha/\alpha \phi(x)$  and  $\alpha/\alpha' f(x)$  with the help of the commutation relations (3 • 12) and (3 • 15),

(3) retain only such terms that do not contain N-product as a factor. aW can reformulate these results by noting that equs.  $(3 \cdot 11)$  and  $(3 \cdot 14)$  can be respectively in the form

$$e^{\pi_{f}}(x) e_{-\pi} = f'(x),$$
  
 $e^{r} \phi^{*}(x) e_{-\pi} = \phi^{*'}(x),$  (3.22)  
 $e^{r} \phi(x) e_{-\pi} = \phi'(x),$ 

where

$$\partial = \frac{1}{2} \int dx dy D(x-y) \frac{1}{2} \frac{1}{2} \int dx dy D(x-y) \frac{1}{2} \frac{1}{2} \frac{1}{2} \frac{1}{2} \int dx dy D(x-y) \frac{1}{2} \frac{1}{2} \frac{1}{2} \frac{1}{2} \frac{1}{2} \int dx dy D(x-y) \frac{1}{2} \frac{$$

as can be readily proved by a direct calculation. Let  $(\phi^*, \phi, S_0)$  be any functional of the :field operators  $\phi^*(x)$ ,  $\phi(x)$  and 'f(x). then

$$(T[\mathscr{O}(\boldsymbol{\phi^{\star}}, \boldsymbol{\phi}, rp)]) = (N[e^{A}e^{A}] (\boldsymbol{\phi^{\star}}, \boldsymbol{\phi}, rp)e^{-A}e^{-E}J). \qquad (3.24)$$

In this formalism the grand partition function given  $\mathbf{b}_y$  (2 • 14) can be put into a compact form:

$$B/\mathcal{B}_{0^{=}} (T[exp-jH_{1}(x)dx]) = (T[\mathbb{R}])$$
$$= (N[e^{a}e^{r} \cdot e e^{-a}e \cdot r])$$

or

$$\Xi/\Xi_0 = \langle N[e^{\Delta}e^{\Sigma}\mathfrak{S}] \rangle, \qquad (3.25)$$

because we can disregard the factor e-ge-::since there is nothing for it to operate on. Serial expansion of exponential function in powers of  $\int H \mathbf{1}(x) dx$  leads us to the fonnula  $(2 \cdot 14b)$ .

In actual calculations dealing with eq.  $(3 \cdot 25)$ , it is more convenient to employ the so-called Feynman graphs. Each tenn in the expansion of the right hand side of  $(3 \cdot 25)$  can be analyzed into various Feynman graphs according to the following rules: On carrying out the rearrangments of operators and retaining only the terms which do not contain. Nproduct, for every factor D(x-x') a dotted (phonon) line is drawn connecting the points x and x'; for every factor S(x-x') a directed (electron) line is drawn from x to x'. Thus each term in the expansion of the right hand side of  $(3 \cdot 25)$  is composed of a number of Feynman graphs, to each of which a product function of S(x-x') and D(x-x') corresponds. The Enal result we want is obtained by integrating with respect to all the coordinates involved in each Feynman graphs and by summing up all the terms contributed from possible Feynman graphs.

#### §4. Dlustrations

Having established the computation rules, we will apply them to the calculations of and C" with small n for the electron-phonon system. What we are going to calculate is

$$= (-1)^{n} (n! J, ... J(T[H1(x_1) ..., H_l(x_n)]) dd ... dx_{x_n},$$

$$H1(x) = g\phi^*(x)\phi(x)rp(x) + g'_{p}(x)_{p}(x).$$
(4.1)

In analyzing \_ into Feynman graphs, the following view points are useful: We think that to each  $\phi^*(x)$  corresponds an electron line starting from the point x, to each  $\phi(x)$ corresponds an electron line entering into the point x, and to each  $r_p(x)$  corresponds a phonon line joining at the point x Thus  $H_1(x)$  represents a point x, at which either three lines. two electron lines and a phonon line, join with strength g, or two phonon lines are connected with strength  $g^\prime$ . We can, therefore, carry out the analysis of  $\_$  by drawing all the graphs in which n vertices are connected with each other, either by two electron lines outgoing and incoming and a phonon line, or by two phonon lines. We need not consider the term in , which contain the factors  $_{rp}(x)$  's of odd numbers. Furthermore, many graphs may be left out of consideration on account of the rule that an electron line is forbidden to join a point to itself. This additional rule comes out from the fact that since a constant factor S(O) corresponds to such an electron line that joins a point to itself, the integration with respect to the coordinate of this vertix is to be reduced to the form jD(x)dx, which evidently vanishes in virtue of  $(3 \cdot 17b)$ . In Fig. 1 various Feynman graphs appearing in the lower order tenns of ,'s are shown. A comparison of the results from Feynman graph analysis with that of a straightfoward application of the computation rules reveals that

the sign of each graph is determined as + or - according to whether the number of closed electron line loops involved in the Feynman graph is odd or even.



Fig. 1. Various Feynman graphs appearing in  $_1$ , 2, 3 and  $_4$ .

Referring to the relations between ... and C" given  $b_y$  (2.16), we can see from Fig. 1 that C" is exclusively constructed of connected Feynman graphs with k vertices. This sesult is of some importance, because the free energy of the system is given  $b_y$ .

$$F = -N\mu - k T \log B$$
  
= -N\mu - k T \log B\_0 - kT (C\_1 + C\_2 + C\_3...), (4.2)

which is to be proportional to the whole volume of the systim V. For every conned Feynman graphs, the result of the integrations with respect to all i;he invJ>lved coordinates but one turns out to be independent of a remaining coordinate, and the integration with respect to the last coordinate simply gives a factor f3V, so that the above requirement that (4.2) is to be proportional to V is always fulfilled if every  $C_k$  consist of connected Feynman graphs alone. It should be noted that this proportionality F ce V is not justified when the averages of Nproducts disregarded above are take into consideration.

The integrations with respect to the coordinates can be quite easily performed, at least for small  $\pi$ . We shall show below only two lowest order terms in Fig. 1

$$CI = \int D(o) dx = f_{3}V \quad iws(Nw+1/2),$$

$$C_{w} = \Pi S(x-x')S(x'-x)D(x-x') dx dx' \qquad (4.3)$$

$$= \left[ -\sum_{k} \sum_{w} \frac{wf_{k}(1-f_{k+w})}{\varepsilon_{k+w}-\varepsilon_{k}+bws} + \sum_{k} \sum_{w} \frac{wN(f_{k+w}-f_{k})}{\varepsilon_{k+w}-\varepsilon_{k}+bws} \right] \beta V,$$

which are in agreement with those obtained by other authors.8>

The merits of the present method are, apart froin the simplicity of its computation rules, that it enables us to get a deep itt&ight into the structures of the higher order perturbations through the Feynman graphs, and hence to go beyond the usual perturbational calculation. For instance, we can carry out a partial summation of serial terms up to infinite order, by adding certain special Feynman graphs. Thus the so-called renormalization procedures developed in quantum field theory will become available to various degrees. An

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• ••

$$\mathbf{O} = (2/2) + (2^2/3) + (2^3/4) + \cdots$$

-С1 + С + О + С  $\cdot \cdot \cdot \cdot \cdot \cdot \bullet =$  a sum ot ан comected Feynman graphs

$$= -0 - (1/2) - (1/4) - (1/2) - (1/4) - (1/4)$$

Fig. 2. Elimination of the term g'rp(x)rp(,c) in H1(,c). This is effected b, employing a re-defined phonon line in place of •••••

example: It is immediately suggested by inspecting Fig. 1 that the effect of the second term in  $(4 \cdot 1)$  can be eliminated by re-defining the phonon line and the phonon energy as illustrated in Fig. 2.

This means in the mathematical formula that D(x-x') is to be replaced by

$$D^{*}(x-x') = D(x-x') - 2g' Jv(x-x_{l}) E(x_{l}-x') dx_{l} + (2g')^{2} \int_{Jv(x-x_{l})} D(x_{l}-x_{l}) D(x_{l}-x_{l}) dx_{l} d$$

The evaluation of  $D^*(x - x')$  is not difficult (see appendix B), the result being conveniently expressed in terms of the Fourier component as

$$D^{*}(k, t) = 1/g' \Gamma' f^{-} D(fk, t) df, \qquad (4.5)$$

where

$$D(k, t) = \int D(x, t) e^{Ik} \cdot dx, \qquad (4.6)$$

and

$$a = (1 + 3g')/g'. \tag{4.6}$$

(4.5) and (4.7) show that for small  $g'D^*(x-x')$  is nearly equal to D(x-x'), whereas for large g' it becomes proportional to 1/g'. This reduction in magnitude of  $D^*(x-x')$ , in turn, acts to prevent the sound velocity from suffering large alteration due to electronphonon interaction. Such a situation remedies a certain difficulty occurred in a perturbational treatment of sound velocity re-normalization.<sup>111</sup> But we shall leave this problem for another occasion.

## § 5. Non-pertu.rbational treatment

The treatment described in the preceding sections is essentially an expansion of the grand partition function in powers of the coupling constant, so that, for the case of strong coupling it will not be useful as it stands. It is, however, possible to put forward a method which is free from the serial expansion procedure.<sup>10</sup>,

We consider a set of functions defined by

$$\begin{aligned}
G_{l}(\mathbf{x}; x') &= (T[\phi^{*}(\mathbf{x})\phi(x')e]) / (T[e]), \\
G_{2}(xy; x'y') &= (T[\phi^{*}(x)\phi^{*}(\mathbf{y})\phi(y')\phi(x')eJ) / (T[e]), \\
G_{3}(xyz; x'y'z') &= (T[\phi^{*}(x)\phi^{*}(y)\phi^{*}(z)\phi(z')\phi(y')\phi(x')\mathbb{R}]) / (T[e]),
\end{aligned}$$
(5.1)

and so on, where

$$\mathfrak{S} = \exp[-\int H_1(x) \, dx], \quad H_1(x) = g\phi^*(x) \, \phi(x) \, p(x). \tag{5.2}$$

(For the sake of simplicity, we shall omit the term g'(f(x)r;(x) for a while.) Following the tenninology used in quantum field theory, we shall call them the Green function of one-electron, of two-electrons and so on. As is evident & om the definitions, however, they correspond to the coordinate representation of the reduced density matrices in the grand canonical ensemble, and play a role similar to the molecular distribution function in classical statistical mechanics. For phonon we define in a similar way

$$\Delta_{I}(xx') = \langle T[\varphi(x)\varphi(x')\mathfrak{S}] \rangle / \langle T[\mathfrak{S}] \rangle, \quad \text{etc.}$$
(5.3)

Now, we will show that these Green functions satisfy a set of coupled integral equations. To do this we apply the equation  $(3 \cdot 24)$  to  $(5 \cdot 1)$  and  $(5 \cdot 3)$ . Then, for instance, we obtain

$$G_{1}(xx') = \langle N[e^{\Delta}e^{\Sigma}\psi^{*}(x)\psi(x')\mathfrak{S}] \rangle / \langle T[\mathfrak{S}] \rangle.$$
(5.4)

Here let us commute  $e^{iJ}$  through  $\varphi^*(x)$ . Referring to the relations

$$e^{J} \Phi^{*} (x) = \Phi^{*} (x) e^{it} + \int_{d_{y}} S(x_{y} - y) a_{y} a_{y} dy e^{it},$$

$$e^{*} \Phi(x) = \Phi(x) e^{iJ} - \int_{d_{y}} S(y - x) a_{y} a dy^{*} (y) e^{it},$$
(5.5)

we see that the result is

$$G_{1}(xx') = \langle N[\epsilon^{\Delta} \int dy S(x-y) \, \delta / \delta \psi(y) \, \epsilon^{\Sigma} \psi(x') \, \mathfrak{S}] \rangle / \langle T[\mathfrak{S}] \rangle. \tag{5.6}$$

The tenn which contains the factor  $\phi^*(x)$  standig to the left of  $e^{it}$  is omitted, because we are  $i_{gn}$  oring the average of Nproduct. Since the operator  $a; a\phi(y)$  commutes with U, we can perform the indicated differentiation in (5.6) by noting that

$$[a/a\phi(y), \phi(x') + = a(_{y} - x'),$$
  
a;  $a\phi(r) = g\phi * (y) (b(y) C,$  (5 -7)

and we obtain

$$G_{I}(xx') = S(x-x') + g f S(x-y) d_{y} (N [e^{A_{e^{U}}} \Phi^{*}(y) \Phi(x') (y) G]) / (T[\mathbb{R}]).$$
(5.8)

Let us further commute  $e^4$  through  $\phi^*(v)$ ,  $\phi(x')$  and  $\phi(v)$ , referring to the relations

$$e^{\Delta}\varphi(y) = \varphi(y)e^{\Delta} + \int dz D(y-z) \,\delta/\delta\varphi(z)e^{\Delta}$$
(5.9)

and

$$a;a(f)(z) = -g\phi^*(z)\Phi(z) C.$$
 (5.10)

The result is

$$G_{1}(xx') = S(x-x') - g^{2} \prod_{d_{y}} \frac{d_{z}}{dz}S(z-y) D(y-z) (T[\phi^{*}(y)\phi(x')\phi^{*}(z)\phi(z)6]) / (T[6])$$

A New Approach to Qualltum-Statistical Mechanics

$$= S(x-x') - g^2 \iint S(x-y) D(y-z) G_2(yz; x'z) dy dz, \qquad (5 \cdot 11)$$

where the use of the definition for  $G_2(\mathcal{W}; \mathbf{x'_y})$  has been made. Thus  $G_1(\mathbf{xx'})$  is shown to be coupled with  $G_2(\mathcal{W}; \mathbf{x'y'})$  through an equation (5 •11). Repeating the same procedures, we can easily prove that  $G_k$  is directly tonnected with  $G_{k-1}$  and  $G_{k+1}$  through an integral equation similar to (5 •11). For example

$$\mathbf{G}_{2}(\mathbf{x}\mathbf{y}; \mathbf{x}'_{\mathbf{y}}') = \mathbf{S}(\mathbf{x} - \mathbf{x}') \quad \mathbf{G}_{1}(\mathbf{y}\mathbf{y}') - \mathbf{S}(\mathbf{x} - \mathbf{y}') \mathbf{G}_{1}(\mathbf{y}\mathbf{x}')$$
$$-\mathbf{g}_{2} \int \mathbf{S}(\mathbf{x} - \mathbf{x}_{l}) \mathbf{D}(\mathbf{x}_{l} - \mathbf{y}_{l}) \mathbf{G}_{s}(\mathbf{y}\mathbf{x}_{l}\mathbf{y}_{l}; \mathbf{y}'\mathbf{x}'\mathbf{y}_{l}) d\mathbf{x}_{l} d\mathbf{y}_{l}.$$
(5.12)

These coupled equations connecting the Green functions of various order bear a resemblance to the integral equations satisfied by the molecular distribution functions of various orders, discovered by Born-Green<sup>12></sup> and Kirdwood<sup>13></sup> in classical statistical mechanics. Although it is a very difficult task to solve these equations, one might be able to find an approximation of breaking off the infinite chain of equations into a closed system of few equations.

It is, however, more convenient to handle an equation containing one electron Green function alone, if such an equation exists. In fact, we can derive such an equation By aking use of a trick of introducing an auxiliary external field.14J We define @ in place of (5 2) by

where  $\phi(x)$  is a c-number field which is to be made vanish in the final result. Then following the same procedures as in deriving (5 •11), we get

$$\mathbf{G}_{1}(\mathbf{x}\mathbf{x}') = \mathbf{S}(, . -\mathbf{x}') \quad \mathbf{I} \quad \mathbf{d}_{\mathbf{y}} \mathbf{S}(\mathbf{x} - \mathbf{y}) (\mathbf{T} \left[ \mathbf{\phi}(\mathbf{x}') \mathbf{\pi} / \mathbf{\pi} \mathbf{\phi}(\mathbf{y}) \mathbf{\Theta} \right]) / (\mathbf{T} \left[ \mathbf{\Theta} \right]).$$

Now in view of (5 •13) it immediately follows that

$$a; a\phi(y) \otimes = \{g\Phi^*(y)q, (v) + \Phi C_y)\Phi^*(y)\} C_{g}$$

and hence

$$G_{1}(xx') = S(x-x') + \mathbf{j} S(x-y) \phi(_{y}) G_{1}(_{y}x') d_{y}$$

$$+ gf d_{y} S(x-y) (N[e^{\pi}e^{\pi}e^{\pi}e^{\Phi} \star (y) \phi(x') < p(y) @]) / (T[@])$$

$$= S(x-x') + \mathbf{j} S(x-_{y}) \phi(_{y}) G_{1}(_{y}x') d_{y}$$

$$- g^{2} \int \int S(x-y) D(y-\hat{s}) G_{2}(y\hat{s}; x'\hat{s}) dy d\hat{s}.$$
(5.14)

The last expression in (5–14) has been attained !: y commuting  $e^{\pi}$  with  $\langle p(y) \rangle$  and carrying out the differentiation. In order to express  $G_2$  in terms of  $G_1$ , we have to regard  $G_1(xx')$ 

as a functional of the auxiliary E.eld  $\phi$  (x) . Then we can proceed as follows :

$$Gi ; {}_{y}f) = (T[\phi^{*}(x)\phi^{*}(f)\phi(f)\phi(y)e]) / (T[e])$$
  
= -(T[\phi^{\*}(x)\phi(y)11/o\phi(f)e]) / (T[e])  
= -af 11\phi(f) (T[\phi^{\*}(x)b(y)e]) / (T[e]). (5 •15)

Inserting  $(5 \cdot 15)$  into  $(5 \cdot 14)$  and carrying out a slight manipulation, we arrive at

$$G(\mathbf{x}\mathbf{x}') = S(\mathbf{x}\mathbf{x}') + \int_{S(\mathbf{x}-\mathbf{y})} \Phi(\mathbf{y}) G(\mathbf{y}, \mathbf{x}') d\mathbf{y} + g^{2} \mathbf{I} \mathbf{S}(\mathbf{x}-\mathbf{y}) D(\mathbf{y}-\mathbf{f}) \mathbf{o}/\mathbf{o} \Phi(\mathbf{f}) G(\mathbf{y}\mathbf{x}') d\mathbf{b} \mathbf{1}_{\mathbf{y}}.$$
 (5.16)

On the other hand, S(x-x') satisfies the following differential equation:

Т

$$\begin{bmatrix} -\underline{H} - 11 & \frac{2}{-\underline{H}} \\ \underline{\pi}t & 2m \end{bmatrix} \mathbf{s}(\mathbf{x}, t) = 11(\mathbf{x}) 11(t) = 11(\mathbf{x}), \qquad (5 \cdot 17)$$

which can be easily proved  $b_y$  a direct operation of  $(\pi/\pi t + \epsilon^2/2\pi \cdot 11)$  on S(x) deE.ned  $b_y$  (3.17a). Operating  $(\pi/\pi t + -b^2/2\pi \cdot \pi)$  on (5.16) from left, and making use of (5.17), (5.16) is converted to

$$\begin{cases} + \frac{11}{2m} = o(\mathbf{x} - \mathbf{x}') + \Phi(\mathbf{x}) G(\mathbf{x} \mathbf{x}') \\ + g^2 \int D(\mathbf{x} - \hat{\boldsymbol{\xi}}) \, \delta / \delta \phi(\hat{\boldsymbol{\xi}}) G(\mathbf{x} \mathbf{x}') d\hat{\boldsymbol{\xi}} . \end{cases}$$
(5.18)

A similar equation for one-nucleon Green function in meson E.eld was solved  $b_y$  Edwards and Peierls  $b_y$  introducing a special technique of Fourier transformation in functional space.<sup>15</sup>> A similar method may be available in the present case.

Another and more tractable method to deal with the grand partition function will be the generalized Hartree approximation. Recently Kinoshita and Nambu have developed a theory of Hartree E.eld for a *system* composed of a number of particles and a intermediar<sub>y</sub> Bose E.eld.<sup>16</sup>> A similar method will be also applicable to the present case, and will be especially useful for the investigation of the cooperative phenomena such as superconductivity.

We shall take again the electron-phonon system as an example. According to the general theory described in § 2 and § 3, the grand partition function is given  $b_{y}$ 

$$\mathbb{E} / \mathbb{E} _{0}^{-} (\mathbb{T} [\exp \left\{-\int \mathbb{H}_{1} (\mathbf{x}) d\mathbf{x}\right\}]), \qquad (5 \cdot 19)$$

which is valid for an arbitrary choice of H  $_0$  and H  $_{1^{\circ}}$  We assume, therefore, that the free Hamiltonian has, instead of (2  $\cdot$ 17b), the following form

$$\bar{H} = \sum_{k} E_{k} a_{k}^{*} a_{k} + \sum_{w} W_{w} (b_{w}^{*} b_{w} + \frac{1}{2})$$
(5.20)

and the interaction Hamiltonian is chosen so as to make it hold that

### A New Approach to Quantum-StatisticaMechanics

$$\int_{HI(x)dx=} \int_{etI70H1e-tii'oJt=\frac{1}{2}H} T[\phi^{*}(x)\phi(x-x')\phi(x')]dxdx' + \frac{1}{4} \int_{I}^{f} P[r(x)x(x-x')r(x')]dxdx' + gj\phi^{*}(x)\phi(x)r(x)dx, \quad (5.21)$$

where  $\phi(x-x')$  represents a sort of Hartee field for electron,  $E_k$  the energy of electron moving in this Hartree field, x(x-x') and Wu, are the corresponding quantities for phonon.  $\phi^*(x)$ ,  $\phi(x)$  and r(x) are respectively defined by

$$\Phi^{*}(\mathbf{x}) = \sum_{k=1/2}^{k} a \mathbf{f} e^{-\mathbf{i}\mathbf{k}\cdot\mathbf{\omega}\cdot\mathbf{E}_{\mathbf{k}}^{\dagger}},$$

$$\psi(\mathbf{x}) = V^{-1/2} \sum_{k} a_{k} e^{i\mathbf{k}\cdot\mathbf{\omega}-\mathbf{E}_{\mathbf{k}}^{\dagger}},$$

$$\varphi(\mathbf{x}) = \sum_{w} (W_{w}/2V)^{1/2} (b_{w}^{*}e^{-i\mathbf{w}\cdot\mathbf{x}+\mathbf{W}_{w}^{\dagger}} + b_{w}e^{i\mathbf{w}\cdot\mathbf{x}-\mathbf{W}_{w}^{\dagger}})$$
(5 •22)

in place of  $(2 \cdot 20)$ . With the aid of  $(5 \cdot 22)$ , each term in the right hand side of  $(5 \cdot 21)$  can be written down as a function of a f,  $a_k$ , b;; and  $b_w$ , the results being

$$H_{[\phi^{*}(x)\phi(x-x')\phi(x')]dxdx'} = \int dt e^{t\vec{x_{0}}} \left[\frac{1}{2} \sum_{k} \phi(k, -E_{k}) (a_{k}^{*}a_{k} - a_{k}a_{k}^{*})\right] e^{-t\vec{H_{0}}},$$

$$H_{4} \int P[r(x)x(x-x')r(x')]dxdx' = \frac{1}{2} \int dte^{t}\vec{u}_{0} \qquad W_{w}x(w, -W_{W}) (bJb_{w} + b_{w}bJ + b_{w}b_{-w} + b_{5}';b_{-w}') Je-tho,$$

$$g \int \phi^{*}(x)\phi(x)r(x)dx = \int dte^{iff_{0}} \left[g_{k} - \frac{(Ww/2V)}{W}\right]^{1/2} (af_{-w}ak_{E}: + ai_{+w}a_{k}b_{w}) Je^{-iff_{0}},$$
(5.23)

where  $\phi(k, E)$  and x(w, W) are, respectively, the Fourier-Laplace transform of  $\phi(x)$  and X(x) defined by

$$\phi(\mathbf{k}, \mathbf{E}) = J \mathbf{j} \Phi(\mathbf{x}, t) e^{\mathbf{i} \mathbf{k}} \mathbf{w} \mathbf{H} J^{\mathbf{B}} \mathbf{x} J \mathbf{t},$$

$$\mathbf{x}(\mathbf{w}, \mathbf{W}) = \prod_{\mathbf{x}(\mathbf{x}, t) \in \mathbf{k}} \mathbf{w} \mathbf{W} J^{\mathbf{3}} \mathbf{x} J \mathbf{t}.$$

$$(5.24)$$

Combining  $(5 \cdot 23)$  with  $(5 \cdot 20)$  we see that

$$\begin{aligned} \mathbf{Fl}_{l} = \mathbf{g} & \underset{\iota}{\text{LJ}} (\mathbf{W}\mathbf{w}/2\mathbf{V})^{1/2} (\mathbf{a}_{k-w}^{*} \mathbf{a}_{w}^{*} \mathbf{b}_{w}^{*} \mathbf{t} + \mathbf{a}_{w}^{*} \mathbf{w} \mathbf{a}_{k}^{*} \mathbf{b}_{w}) + \underset{k}{\Phi} (\mathbf{k}, -\mathbf{E}_{k}) \text{ af } \mathbf{a}_{k-\frac{1}{2}} & \underset{k}{\Phi} (\mathbf{k}, -\mathbf{E}_{k}) \\ & + \frac{1}{2} \sum_{w} \mathcal{W}_{w} \chi (w, -\mathcal{W}_{w}) (b_{w}^{*} b_{w} + b_{w} b_{w}^{*} + b_{w}^{*} b_{-w} + b_{w}^{*} b_{-w}^{*}). \end{aligned}$$

Since the sum of HD and  $H_1$  has to be taken equal to the original total Hamiltonian given

by (2.17), apart from a constant,

$$H0+H = H+C$$

it follows by comparing  $(5 \cdot 25)$  with  $(2 \cdot 17)$  that

$$S_k = E_{\bar{k}} + \phi(k, -E_k),$$
 (5.26a)

$$\hbar w s = W_w,$$
 (5 •26b)

$$g' = x(w, -Ww),$$
 (5.26c)

$$C = -\frac{1}{2} 2 J \Phi(k, -E_k).$$
 (5.26d)

Eqs. (5.26) provide physical meanings for the Hartree field  $\phi(x)$  and x(x), that is, the Fourier-Laplace component of  $\phi(x)$  equals the difference between the enenely cf a free electron and that of an electron moving in the Hartree field, and the Fourier-Laplace component of X(x) gives what we called renormalization constant in § 2. In order to determine  $\phi(x)$  and x(x) self-consistently, we shall apply the theory of Green function described in the beginning of this section. Define the Green function G(xx') and LI(xx') by

with

$$e = e \times p[- \{g] \Phi^{*}(x) \phi(x) So(x) dx + JJ \phi^{*}(x) \phi(x-x') \phi(x') dx dx' + \frac{1}{4} H_{3}(x) \times (x-x') g(x') dx dx'\}].$$
(5.28)

If we content ourselves with the expressions up to the second order in the coupling constant g, the coupled equations for G(x-x') and LI(x-x'), which are derived through the procedures described above, can be easily solved by an iteration procedure. We shall give here only the results;

$$G(xx') = S(x-x') + \frac{1}{2} H S(z-y) \phi(x-y) S(z-x') d_y dz$$
  
+  $g^2 \iint_{S(x-y)D(y-z) S(y-z) S(z-x') dy dz},$  (5.29)

$$g(xx') = D(x-x') - \frac{1}{2} H D(x-y) x(y - z) D(z - x') dy dz$$
  
$$+ g^{2} \iint D(x-y) D(x'-z) S(y - z) S(z - y) d_{y} dz.$$
(5.30)

In this approximation, it will be natural to determine  $\phi(x)$  and X(x) in such a way that

$$G(xx') = S(x-x'), \quad LI(xx') = D(x-x'), \quad (5.31)$$

because the equations (5  $\cdot$ 21) state that electrons and phonons behave in the respective

Hartree field as if they are  $ind_{ep}$  endent of each other. This statement is in accord with the bask assumptioninHartreeapproximation. From (5.29), (5.30) and (5.31)  $\phi(x)$  and x(x) are determined as

$$\phi(z-y) = -2g2D(y-z)S(y-z), \qquad (5 \cdot 32)$$

$$x(y-z) = -2g^{2}S(y-z)S(z-y)$$
 (5.33)

Here we shall briefly discuss the results derived from the equations  $(5 \cdot 32)$  and  $(5 \cdot 33)$ , leaving the details to a later publication.

Performing the Fourier Lapace transformations of  $(5 \cdot 32)$  and  $(5 \cdot 33)$  with the help of  $(3 \cdot 17)$ , we readily get

$$\phi(\mathbf{k}, -E_{k}) = \frac{g^{2}}{V} \sum_{w} \hbar w s \left[ \frac{(N_{w}+1)(1-f_{k-w})}{E_{k-w}-F_{k}+\hbar w s} + \frac{N_{w}(1-f_{k+w})}{E_{k+w}-E_{k}-\hbar w s} \right], \quad (5.34)$$
$$\mathbf{x}(\mathbf{w}, -\mathbf{W}_{k}) = \frac{2g^{2}}{V^{k}} \frac{[\mathbf{k}(1-f\mathbf{k}-\mathbf{w})]}{\mathbf{E}_{k}-\mathbf{w}-\mathbf{E}_{k}+\mathbf{W}\mathbf{w}}. \quad (5.35)$$

Referring to  $(5 \cdot 26)$ , the equation  $(5 \cdot 35)$  gives the change in the sound velocity of phonon caued by the electron phonon interaction, if the dispersion of sound velocity is ignored. The result obtained here is nearly the same as that calculated by Frohlich.  $(5 \cdot 34)$  combined with  $(5 \cdot 26a)$ , on the other hand, give an equation to determine  $\overline{E}_k \cdot On$  neglecting N w at very low temperatures, it becomes

$$E_{k} = \ddot{e}_{k} - L \qquad 8_{w} s \frac{(1 - fk - u)}{E_{k} - v} - E_{k} + b_{w} s}. \qquad (5.36)$$

It is interesting to note that the same equation as (5.36) was derived by Bardeen in a quite different way.17> Although the nature of the solution of the equation (5.36) was already discussed by him, a more careful investigation of this equation has been made by the present author. The result obtained agrees with that given by Bardeen in its essential point. There exists a solution of (5.36) in which one electron energy E, has a gap at Fermi surface for sufliciently strong coupling constant. The ground state in which all the states of lower energy are occupied by electrons will correspond to a superconducting state at O'K. Temperature effect on energy spectrum is easily taken into account in the present method, and it is shown that the energy gap, dependent on temperature, becomes to vanish above a certain temperature. Thus a sort of phase transition is expected. A similar result was worked out by Frohlich and Kuper<sup>18 J</sup> with one-dimensional model. The basic idea of the present method resembles rather that of Frohlich's than Bardeen's. Frohlich has assumed that a cooperative interaction between electrons and phonons produces such a potential for an electron as to give rise to a splitting of the energy spectrum of the electron. In the present theory, we introduced explicitly a possessing a nature which Frohlich has assumed, and we have proved in a self-consistent manner that this potential actually gives rise to a splitting in one-electron spectrum even in three dimensional case, and hence brings the assembly of electrons into a special state which we want to call the superconducting state.

# §6. Extension to Other Systems

The electron-phonon system so far considered. is a typical example, to which our method can conveniently be applied. In extending our method to other system, it may happen that some modifications in the formalism are required. In this section we shall show that our method is easily extended, to a system in which many particles, obeying Fermi or Bose statistics, are interacting with each other through two-body potential.

In the scheme of second quantization we write the Hamiltonian of a systen in question as

$$H^{-} H_{0} + H_{1},$$

$$H_{0} - \int \phi^{*}(x)p^{2}/2\tau \phi(x) d^{3}x,$$

$$H_{1} = \frac{1}{2} \int \phi^{*}(x)\phi^{*}(x')J(x-x')\phi(x')\phi(x) d^{3}x d^{3}x^{4},$$
(6.1)

where  $\int (x - x^{l})$  represents the interaction potential between two particles located at x and x', p the momentum operator of a particle.  $\phi^{*}(x)$  and  $\phi(x)$  can be expanded. Into Fourier series :

$$\phi^{*}(x) = \frac{v - 1/2}{k}$$
 aj;e-ik····,  $\psi(x) = V^{-1/2} \sum_{k} a_{k} e^{ik \cdot x}$ , (6·2)

in which aj; and  $a_{1,}$  are as usual creation and annihilation operators of a particle with wave vector  $k_{c}$  the commutation relations between them being given by

The grand partition function 8 can be written as

$$8/8 \circ 1 + 1 + 2 + 3 + \cdots$$
  
= exp(C<sub>1</sub> + C<sub>2</sub> + C<sub>3</sub>...) (6.4)

with

$$\boldsymbol{\xi}_{n} = (-1)^{n} / n! \int \cdots \int \langle P[H_{1}(t_{1}) \cdots H_{1}(t_{n})] \rangle dt_{1} \cdots dt_{n},$$

$$H_{1}(t) = \boldsymbol{e}^{tH_{0}} H_{1} \boldsymbol{e}^{-tH_{0}},$$
(6.5)

or more formally as

$$\mathbb{Z}/\mathbb{Z}_{0} = \langle P[\mathfrak{S}] \rangle,$$

where

$$\mathfrak{S} = \exp[-\int H_1(t)dt].$$

First we consider the case of Fermi particles. In applying the computation rules derivi in § 3 to the present case, a difficulty arises from the interaction term

A New Appmoach to. Quantum-Statistical Mechanics 371

$$\int H_1(t) dt = \frac{1}{2} \iint \psi^*(x, t) \psi^*(x', t) J(x - x') \psi(x', t) \psi(x, t) d^3x d^3x' dt, \qquad (6.7)$$

which prevents us from accomplishing in a simple manner the t-ordering indicated in the right hand side of (6.5). The problem can, however, be solved  $B_y$  rewriting (6.7) in the form

in which ] (x - x') is defined  $\mathbf{b}_{v}$ 

-

That (6.8) equals (6.7) is easily verified  $b_y$  a short calculation. Then, noting that for arbitrary T-products T[A], T[B],...

we obtain, in place of (6.6),

with

$$\Xi/\Xi_0 = \langle T[\mathfrak{S}] \rangle$$

$$e = e \times p [-\frac{1}{2} \iint_{\varphi} (x) \varphi(x)](x - x') \varphi^*(x') \varphi(x') dx dx'$$
(6.10)

to which all the rules established in § 3 are now applicable.

We have no trouble with the case of Bose particles. We do not want to repeat here cslong analysis, so that we give below only the lemma II modified so as to hold for both statistics.

Lemma III For any product of  $\phi^*(x)$  and  $\phi(x)$  denoted by  $(\phi^*, \phi)$ , it holds that

$$T[\mathfrak{H}(\psi^*,\psi)] = N[\mathfrak{H}(\psi^{*\prime},\psi^{\prime})],$$

in which T = (=F1)<sup>P</sup> P and  $\phi'(x)$  and  $\phi'(x)$  are defined  $\mathbf{b}_{v}$ 

$$\phi^{*'}(x) = \Phi^{*}(x) + jdx'S(x-x')d/d\phi(x'),$$

$$\phi'(x) = \phi(x) = jdx'S(x'-x)d/d\phi^{*}(x'),$$

respectively.  $d/d\phi^*(x)$  and  $d/d\phi(x)$  satisfy the following commution relations:

$$[ \underline{A} / \underline{A} \phi(\mathbf{x}), \phi(\mathbf{x}') ]_{\pm} = [ \underline{A} / \underline{A} \phi^{*}(\mathbf{x}), \phi^{*}(\mathbf{x}') ]_{\pm} = \underline{A} (\mathbf{x} - \mathbf{x}'),$$
$$[ \underline{A} / \underline{A} \phi(\mathbf{x}), \phi^{*}(\mathbf{x}') ]_{\pm} = [ \underline{A} / \underline{A} \phi^{*}(\mathbf{x}), \phi(\mathbf{x}') ]_{\pm} = \mathbf{O}.$$

S(x-x') is given by

$$S(x-x') = \begin{cases} V^{-1} \sum_{k} f_{k} e^{-ik \cdot (x-x') + \varepsilon_{k}(t-t')} & t > t' \\ V^{-1} \sum_{k} (f_{k} \mp 1) e^{-ik \cdot (x-x') + \varepsilon_{k}(t-t')} & t < t' \end{cases}$$

where

 $f_k = (e^{\alpha + \beta \varepsilon_k} \pm 1)^{-1}.$ 



Fig. 3. Various Feynman graphs appearing in C1 and C2 Dotted line  $x \cdots x'$  corresponds to j(x-x'). Directed line  $x \cdot xI$ corresponds to S(x-x). X represents self energy ](O).

All the upper  $si_{gn}s$  ot doubled  $si_{gn}atures$  correspond to Fenni case and lower  $si_{gns}$  to Bose case.

W e can analyze or ck into Feynm.an graphs drawing a directed Ь, (particle) line for every factor S(x-x') from x to ,!, and a dotted line connecting the points x and ,! for every factor ] (x - x'). In the present case a particle line may join a point to itself. In Fig. 3 the Feynman graphs appearing in the lower order tenns of  $C_{k}$ 's are shown. It will be worth while to compare the results

derived from Fig. 3 with

that of Ichimura's calculation. In the notation adopted here, the results for  $C_{k'}$ s obtained by Ichimura  $\mathbf{b}_{v}$  his own method are written as follows :<sup>8></sup>

$$C_{1} = -\frac{1}{2} \{ 1V - 1 \sum_{k \neq J} 2J(]o'=F]k_{-i} \}_{k}^{f} \{ , \qquad (6 \cdot ||a) \}$$

$$C_{2} = \frac{1}{2} \frac{1}{2} J 2J_{2} J 2J_{2} J \frac{1}{k} (1 \neq \{k\} \text{ fifm} Uo=F]k_{-1}) Uo=F]k_{-T} \}$$

$$+\frac{1}{4} \{ 1V - \frac{2}{2} J 2J_{2} J 2J_{2} J \frac{1}{k} (J \neq \{k\} \text{ fifm} Uo=F]k_{-1})^{2} (1 \neq \{k\} (1 \neq \{k\}) \text{ fifn}, \frac{k_{k}}{k_{-1}} + \frac{1}{2} \cdot ck + si - Em - cfl \} \}$$

$$(6 \cdot ||b|)$$

where  $\mathbf{j}_{\mathbf{k}}$  is the Fourier component of f(x) defined  $\mathbf{b}_{\mathbf{v}}$ 

$$]_{k} = jJ(x)e1^{k} \cdot OO^{8}x.$$

We will show that (6.11) are in complete agreement with our results. Inspecting Fig. 3,

our expression for  $C_{\!\!1}$  is readily written dQwn as

$$C_{1} = \frac{1}{2}(0) \int_{S(0)} dx - \frac{1}{2}H \int (x_{1} - x_{2}) \{S(0)\} \frac{1}{2dxiJx^{2}} \\ \pm \frac{1}{2} \int \int (x_{1} - x_{2}) S(x_{1} - x_{2}) S(x_{2} - x_{1}) dx_{1} dx^{2}.$$
(6.12)

Noting that

$$S(o) = v - l \, \mathop{}^{"i:ifk}_{k}$$

$$S(x - , l) \, S(, l - x) = -v - \ddot{o} \, \mathop{k}^{fk}(l = \mathcal{H}[k_l] \, e'(k - k') \cdot (\mathcal{H}e^{-jk}) - (\mathcal{H}e^{-jk}) \, l, -, l, l, k = \mathcal{H}[k_l] \, e'(k - k') \cdot (\mathcal{H}e^{-jk}) - (\mathcal{H}e^{-jk}) \, l, -, l, k = \mathcal{H}e^{-jk} \, dk = \mathcal{H}e^{-jk$$

(6 • 12) can be reduced to

$$Cif.BV = -\frac{1}{2}V^{-2} \{ \{ -2\}k[k, \pm lofk[k, \pm f]kl - k\{k([k, \pm f])\} \}$$
$$= -\frac{1}{2}V^{-2}\sum_{k}\sum_{k'} (J_0 \mp J_{k'-k})f_k f_{k'},$$

which is just equal to  $(6 \cdot 11a)$ . It is seen from the above calculation that the term  $J(O) J\Phi^*(x)\phi(x)dx$  in @250 behaves as to subtract the self-energy from the final results. We *may*, therefore, disregard this term hereafter, provide we keep in mind that the self-energy parts are always to be subtracted. Then  $C_2$  is shown to consist of five integrals:

$$C_{2} = =F4C_{M} + 2C_{2}s + BC_{2}0 = F4C_{2}D = F2C_{2}E_{1}$$

$$C_{M} = \frac{1}{4}\int_{0}^{\infty} \frac{1}{1}\int_{0}^{\infty} \frac{1}{2}J(x_{1} - x_{2})S(x_{2} - x_{3})S(x_{3} - x_{2})J(x_{8} - x_{4}) dd \cdots dx4$$

$$=\frac{1}{4}\int_{0}^{2}V^{-2}J - \frac{1}{k}\int_{0}^{\infty} \frac{1}{k}\int$$

As to the evaluation of the above integrals, see appendix B. The summation of these five integrals with given numerical coefficients immediately leads us to (6·IIb), apart from the self-energy parts which can be cancelled out by the integrals corresponding to the graphs  $C'_{2.4}$ , C'<sub>2</sub>B and C'<sub>2</sub>o in Fig. 3.

An application of the method of partial summation over certain special Feynman graphs, which was proved useful in § 4, gives rise to an interesting result for the electrons interacting with coulomb potential. It is suggested from Feynman graph analysis that the interaction potential ](x - x') has better to be replaced by

$$\int [f(x-x') = \int (x-x') + 2 H J (x-x_1) S(x_1 - x_2) S(x_2 - x_1) J (x_2 - x') dx dx_2 + 2^2 \int [f] f \int (x-x_1) S(x_1 - x_2) S(x_2 - x_1) \int (x_2 - x_3) S(x_3 - x_4) S(x_4 - x_3) \int (x_3 - x') ax' Qx_4 + ''' (6 \cdot 13)$$

For coulomb potential  $](x - x') = e^{2t} |x - x'| \mu(t - t')$ , the Fourier transform of (6-13), integrated over t, becomes

$$J^{*}(\mathbf{k}) = \frac{4\pi e^{2}}{k^{2}} \left\{ 1 + \frac{4\pi e^{2}}{k^{2}} \Lambda(\mathbf{k}) + \left(\frac{4\pi e^{2}}{k^{2}} \Lambda(\mathbf{k})\right)^{2} + \cdots \right\}$$
  
=  $\frac{4 \sin^{2}}{k^{2} - 4 \cdot \sin^{2} \Pi(\mathbf{k})}$   
$$\pi c k^{2} = \sum_{V = kl} \sum_{kl=1}^{l} s(k'; -t) S(k' - k; t) at = \sum_{V = kl} \sum_{kl=1}^{l} \frac{f(\mathbf{k}, -f(\mathbf{k}, -\mathbf{k}))}{s_{k}, -s_{k} \Gamma_{k}} \cdot (6 \cdot 14)$$

This result shows us that the coulomb interaction between electrons is to be screened (as 1/r-e-J.r/r), the screening constant A being roughly estimated as

$$;_{2}=_{4}:ire^{2}\Pi(0) = -\frac{8ire^{2}}{V}: \underset{k}{E} \left(\frac{3fk}{\partial B_{k}} - \frac{4ire^{2}m}{B^{2}}K,... \right)$$
(6.15)

(K,,.=the magnitude of wave vector of electron with Fermi energy).

The same result as (6 • 15) for the screening effect was derived by Macke by a variational calculation.19>

# Appendix A

## A Proof of Lemma 1

We shall prove the validity of lemma I by a mathematical induction. Assume that

it is tme for some T-product TIIJ(So)], i.e. that

$$\Pi \amalg (So) = N [g: (So')]. \tag{A.1}$$

Then if we can prove it for  $T [S_0(x) g: (S')]$ , we can conclude that the statement of lemma I is valid, since for  $g:(S_0)=1$  and for  $g:(S_0)=S_0(A \cdot 1)$  is trivially true. Without a loss of generality we can take  $g:(S_0)$  as a product of  $\pi$  factors;  $S_0(x_1) S_0(x_2) \cdots S_0(x_n) \cdot When$ 

$$t_1 > t_2 > \cdots > t_n$$

ТПJ(So)] can be written as

$$T[\mathfrak{F}(\varphi)] = \varphi(x_1)\varphi(x_2)\cdots\varphi(x_n). \qquad (A\cdot 2)$$

Now let us assume that

$$t_1 > t_2 > \cdots + t_r > t > t_{r-1} > \cdots > t_n$$
.

Then

$$T[S_0(x) g: (S_0)] = S'(x_1) \cdots S_0(x_r) S_0(x) S_0(Xr+1) " \cdot S_0(x_{m-1})$$
  
=X(S\_0)S\_0(x) Y(S\_0)- (A · 3)

Here we decompose  $S_0(x)$  into two parts according to

$$S_0(x) = S_{0+}(x) + S_{0-}(x)$$

and transfer S<sub>0</sub>-(x) to the left through  $X(S_0)$  and  $S_0+(x)$  to the right through  $V(S_0)$ . The result is conveniently expressed as

$$X(\varphi)\varphi(x)Y(\varphi) = \varphi_{-}(x)XY + XY\varphi_{+}(x) + \int dx' \mathcal{A}_{1}(xx')\delta/\delta\varphi(x')(XY) \quad (A\cdot4)$$

where

In order to obtain the N-product of  $X_{S_{1}}(x) \times from (A \cdot 4)$ , we have to bring back  $s_{1}(x)$  to the right and  $S_{0^{+}}(x)$  to the left, because  $S_{0^{-}}(x)$  may stand to the left of  $S_{0^{-}}(x_{1})$ ,  $S_{0^{-}}(x_{2})$ ,  $\cdots S_{0^{+}}(x_{1})$ , and  $S_{0^{+}}(x)$  may stand to the right of  $S_{0^{+}}(X_{1+1})$ ,  $\cdots S_{0^{+}}(x_{1-1})$ . Carrying out rearrangements needed for getting N-product, we can express the result in the form

$$So_{-}(x)XY = N [So_{-}(x)XY] + jdx'[So_{-}(x), So_{-}(x')]_{i}3X/i3So_{-}(x') Y,$$

$$XY So_{+}(x) = N [X + Y So_{+}(x) J - jdx'[So_{+}(x), So_{+}(x')]_{x}^{x}BY / BSo_{+}(x'),$$
(A.6)

where operators i3/i3So+ (x) and i3/i3So (x) are difined through

$$[i3/i3S_{0+}(x), S_{0+}(x')] = [i3/i3S_{0-}(x), S_{0-}(x')] = \partial (x - x'),$$
  
$$[i3/i3S_{0+}(x), S_{0-}(x')] = [i3/i3S_{0+}(x), S_{0+}(x')] = 0,$$
  
(A.7)

namely they represent functional differentiation. Since X and y contain  $P_+(x)$  and  $P_-(x)$  as factors through the combined form  $f(x) = f(+(x) + P_-(x))$ , it follows that

$$iJX/iJ'P-(x) = iJX/iJrp(x), \quad iJY/iJ'P+(x) = iJY/iJ(f)(x).$$
 (A-8)

If we define further

$$\partial_{2'}(xx') = \{ \begin{array}{cc} [\cdot\mathbf{P} + (x)' \ \mathbf{P} + (x')]_{-1} & t > t' \\ -[\mathbf{r}\mathbf{p}_{-}(\mathbf{x}), \ \mathbf{P} - (\mathbf{x}')]_{-1} & t < t', \end{array}$$
(A·8)

then from  $(A \cdot 3) - (A \cdot 8)$  we obtain

$$X(f)(x) Y = N [ {P+ (x) + rp_(x)} X Y] + Jdx' [1](xx') - L 1_2(xx')] g (X Y) / g(J)(x').$$
  
(A·IO)

In view of the first assumption (A 1) and a property of N-product

$$N[\varphi'N[\mathfrak{F}(\varphi')]] = N[\varphi'\mathfrak{F}(\varphi')]$$

(A 10) tells us that

$$T[\varphi(x)\mathfrak{F}(\varphi)] = N[\varphi'(x)\mathfrak{F}(\varphi')]$$

where

$$f(x) = (f)(x) + \int dx'_{\{1_1(xx') - 1i xx'\}} \frac{8}{4} g(J)(x')$$

With the help of (A-5) and (A-9) it is easy to show that

$$\frac{\partial I(\mathbf{x}\mathbf{x}')}{\langle \mathbf{f}_{+}(\mathbf{x}), (\mathbf{f})(\mathbf{x}') \rangle - [\mathbf{f}\mathbf{f}\mathbf{J}_{+}(\mathbf{x}), \mathbf{f}\mathbf{J}_{+}(\mathbf{x}')] - [\mathbf{f}^{-}\mathbf{f}^$$

which is identical with the definition of D(x - x') given by (3.8). Thus our proof is completed. The proof of lemma II cn be achieved in quite a similar manner, so that it will be unnecessary to repeat here the similar procedure.

### Appendix **B**

First we shall derive the formula  $(4 \cdot 5)$ . For simplicity, we employ hereafter such an unit as to make s = 1 and  $'\overline{o} = 1$ . Fourier component of D(x) is then expressed as

$$D(k, t) = \frac{1}{2}k\{(N_k+1)e^{-k|t|} + N_k e^{k|t|}\}.$$
 (B 1)

Now the Fourier transform of (4-4) is given by

 $D^{*}(k, t-t') = D(k, t-t') - 2g^{-1}f : D(k, t-t_{-1})D(k, t_{1}-t') dt$ 

.d New Approach to Quantum-Statistical Mechanics

+ 
$$(2g^{1})^{2}J: J > (k, t-t_{1})D(k, t_{1}-t_{2})D(k, t_{2}-t')dt_{1}dt_{2}''$$
 (B·2)

Although  $D^*(k, t)$  can be evaluated by solving an integral equation

$$D^{*}(k, t) = D(k, t) - 2g'f:D(k, t-s)D^{*}(k, s)ds,$$
 (B-3)

we shall follow a more direct and elementary method. A short calculation yields that

$$\int_{0}^{\beta} D(k, t-t_{1}) D(k', t_{1}-t') dt_{1} = \frac{k^{2}}{k^{2}-k'^{2}} D(k, t-t') - \frac{k'^{2}}{k'^{2}-k^{2}} D(k', t-t') \quad (B\cdot4)$$

which is reduced in the limit of k-+k' to

$$\int_{0}^{\beta} D(\mathbf{k}, t-t_{1}) D(\mathbf{k}, t_{1}-t') dt_{1} = (1-\frac{1}{2}kd/dk) D(\mathbf{k}, t-t') = \mathcal{A}_{k} D(\mathbf{k}, t-t') \qquad (B\cdot 5)$$

where

$$dk = (1 - \frac{1}{2}kd/dk).$$
 (B·6)

Using (B•5) in a repeated manner; we can put (B•2) into the form

$$D^*(\mathbf{k}, t) = (1 - 2 \mathbf{g}^{\mathsf{T}} \mathbf{d}_{\mathsf{t}} + (2 \mathbf{g}^{\mathsf{T}} \mathbf{J}_{4-}^{\mathsf{T}} - \cdots) D(\mathbf{k}, t)$$
$$-\frac{1}{1 + 2 \mathbf{g}^{\mathsf{T}} \mathbf{J}_{2}} D(k, t),$$

or

$$(1+2g'\mathcal{A}_k)D^*(\boldsymbol{k},\,t)=D(\boldsymbol{k},\,t). \tag{B.7}$$

It is easy to solve this inhomogeneous linear differential equation of first order with a condition  $\lim_{k \to m} D^*(k, t) = 0$ . The result is

$$D^{*}(k, t) = 1/g'[k^{H} D(k', t)k'-"dk'$$
  
= 1/g'f:f-"D(fk, t)d (B.8)

where

$$a = (1 + 3g')/g'.$$

Thus (4.5) is proved. For the Fourier component of S(x), the following equations are easily proved :

$$\int_{0}^{\beta} S(k, t-t_{1}) S(k', t_{1}-t') dt_{1} = \frac{S(k, t-t') - S(k', t-t')}{\varepsilon_{k} - \varepsilon_{k'}}, \quad (B.9)$$

$$1:S(k, t-t_{1}) S(l, t_{1}-t) S(m, t_{1}-t') S(n, t'-t_{1}) dt_{1}$$

$$= (f_{m,-} \{ , . ) / (8_{i} + 8_{m,-} - 8_{k} - 8_{n}) S(k, t-t') S(l, t'-t) \quad (B.10)$$

$$(fk - fi)/(ei + em - e_k - E_{\Pi})S(m, t - t')S(n, t' - t).$$

In particular, in the limit of  $k \rightarrow k^{T}$  tt' (B.9) becomes

$$\int_{0}^{r} S(\mathbf{k}, \mathbf{t} - \mathbf{t}_{i}) S(\mathbf{k}, \mathbf{t}_{i} - \mathbf{t}) d\mathbf{\hat{\mathbf{t}}} = -S(\mathbf{k}, \mathbf{o}) = -\mu \mathbf{f}_{k} (1 = \mathbf{F}[\mathbf{\hat{k}}] - (\mathbf{B} \cdot \mathbf{1})$$

(B•9), (B•10) and (B•11) were used in evaluating the integrals  $C_{2A}$ -  $C_{2lii}$  in §6.

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