## 4 Diagram notation

**Notation 4.1.** *Diagram notation.* In diagram notation, particle-hole operators are written as oriented lines extending from a vertex. Particle annihilation operators enter the vertex from below, particle creation operators leave the vertex at the top, and single-excitation operators have both creation and annihilation lines. Contractions are represented by joining particle-hole lines with compatible position and orientation.

$$\stackrel{\uparrow}{\uparrow} \equiv a_p \qquad \qquad \stackrel{\downarrow}{\uparrow} \equiv a_p^{\dagger} a_q = a_q^p \qquad \qquad \stackrel{\uparrow}{\uparrow} \equiv \overline{a_p} \overline{a_q}^{\dagger} = a_p^{q^{\bullet}} \qquad (4.1)$$

Quasiparticle operators are distinguished by the use of closed-circle vertices, with particle lines pointing upward and with hole lines pointing downward. Single-excitation operators split into four cases representing the virtual and occupied blocks of  $a_a^p$ . Internal contractions of single excitations (bubble contractions) are implicitly taken to be hole contractions.

Higher excitation operators are depicted by joining single-excitation operators with a solid line. Contracted operators are implicitly normal ordered together. Normal-ordered products of uncontracted operators are joined with a dotted line.

$$\underbrace{\underbrace{-\cdots +}_{m \text{ times}}}_{\text{times}} \equiv :a_{q_1}^{p_1} \cdots a_{q_m}^{p_m} := a_{q_1 \cdots q_m}^{p_1 \cdots p_m}$$

$$= :a_{q_1 \cdots q_m}^{p_1 \cdots p_m} a_{s_1 \cdots s_n}^{r_1 \cdots r_n} :$$

$$\underbrace{-\cdots +}_{m \text{ times}} = :a_{q_1 \cdots q_m}^{p_1 \cdots p_m} a_{s_1 \cdots s_n}^{r_1 \cdots r_n} :$$

$$\underbrace{-\cdots +}_{m \text{ times}} = :a_{q_1 \cdots q_m}^{p_1 \cdots p_m} a_{s_1 \cdots s_n}^{r_1 \cdots r_n} :$$

 $\Phi$ -normal-ordering is indicated by the use of double-circle vertices, @ and @ instead of o and  $\bullet$ . These building blocks can be used to construct more complicated products. Products of diagrams are written from top to bottom, representing left-to-right ordering in the corresponding algebraic expression.

**Definition 4.1.** m-electron operators in diagram notation. The building blocks of a graph are m-electron operators, which can be represented in two equivalent ways. The Goldstone representation depicts an operator as a label attached to the corresponding excitation operator, whereas the Hugenholtz representation depicts the operator as a single vertex with m outgoing and incoming lines. Note that  $\left(\frac{1}{m!}\right)^2 \sum_{\text{Einstein}}$  is baked into the definition (see def 4.2 and axiom 4.1).

$$\boxed{\boldsymbol{v}} \xrightarrow{\boldsymbol{\psi}} \cdots \xrightarrow{\boldsymbol{\psi}} \equiv \left(\frac{1}{m!}\right)^2 \sum_{\text{Einstein}} \overline{v}_{p_1 \cdots p_m}^{q_1 \cdots q_m} a_{q_1 \cdots q_m}^{p_1 \cdots p_m} = \overbrace{\boldsymbol{v}}^{\boldsymbol{v}} \underbrace{\boldsymbol{v}} \cdots \xrightarrow{\boldsymbol{v}} = \overline{v}_{p_1 \cdots p_m}^{q_1 \cdots q_m} a_{q_1 \cdots q_m}^{p_1 \cdots p_m} a_{q_1 \cdots q_m}^{q_1 \cdots q_m}$$

$$(4.4)$$

$$\underbrace{v} \longrightarrow \underbrace{\left(\frac{1}{m!}\right)^2}_{\text{Einstein}} \underbrace{\sum_{\text{Pintein}}}_{p_1 \cdots p_m} \underbrace{\overline{a}_{q_1 \cdots q_m}^{p_1 \cdots p_m}}_{p_1 \cdots p_m} \underbrace{v}_{q_1 \cdots q_m} = \underbrace{v}_{q_1 \cdots q_m} \underbrace{v}_{q_1 \cdots q_m} \underbrace{v}_{q_1 \cdots q_m} \underbrace{v}_{q_m} \underbrace{v}_{q_1 \cdots q_m} \underbrace{v}_{q_2 \cdots q_m} \underbrace{v}_{q_2$$

The labeled diagrams on the right represent just the summand of the operator, which highlights the difference between representations. Both summands correspond to an excitation operator weighted by its antisymmetrized interaction tensor, but whereas the Goldstone summand specifies an ordering for the indices of its corresponding algebraic term, the Hugenholtz summand does not. Since the phases of  $\overline{v}_{p_1...p_m}^{q_1...q_m}$  and  $a_{q_1...q_m}^{p_1...p_m}$  cancel under index permutation, the two labeled diagrams are actually equal – a Hugenholtz summand can be expanded into a Goldstone summand by simply choosing an arbitrary ordering for the indices. In practice, the symmetry of the Hugenholtz operator simplifies the enumeration of Wick expansions whereas the Goldstone operator makes it easier to evaluate a graph's overall phase.

<sup>&</sup>lt;sup>1</sup>In the original paper [J. Goldstone, *P. Roy. Soc. A* **239**, (1957)], Goldstone's diagrams were actually defined in terms of non-antisymmetrized integrals. The *antisymmetrized Goldstone diagrams* used here are sometimes called *Brandow diagrams*.

**Definition 4.2.** Graph. A graph<sup>2</sup> G = (O, L, h, t) consists of a set of m-electron operators O, a set of lines L, and two mappings  $h, t: L \to O$  that return the head  $h(l) \in O$  and tail end  $t(l) \in O$  of each line in L. Two lines are considered equivalent if they share the same head and tail. Here, we allow for external lines in which one of the ends is e, the free end, which is formally considered a member of O. Lines with no free end are termed internal. A graph is termed closed if it contains no external lines, representing a scalar-valued algebraic term. Otherwise, the graph is open and represents an operator-valued algebraic term. The rules of interpretation for translating G into an algebraic expression are given below.

**Definition 4.3.** Connected and linked graphs. Two lines are adjacent if they share a non-free end. In a Goldstone diagram, two lines are termed Goldstone adjacent if they end on the same single-excitation vertex of an operator. In this context, the broader sense of the adjacency is called Hugenholtz adjacency. A path is a sequence of lines  $(l_1, \ldots, l_n)$  such that  $l_i$  is adjacent to  $l_{i+1}$  and no line is repeated. In a Goldstone diagram, one can distinguish between Goldstone and Hugenholtz paths depending on whether or not the lines are all Goldstone adjacent. A Goldstone cycle is a Goldstone path whose ends are either Goldstone adjacent or free. A Goldstone cycle with free ends is an open cycle. Otherwise, it forms a loop. A graph is considered connected if there is a Hugenholtz path connecting any two of its operators. A closed, disconnected subgraph is considered unlinked from the rest of the graph. Note this last distinction: A disconnected graph is still classified as linked, as long as none of its disconnected subgraphs are closed.

**Definition 4.4.** Equivalent subgraphs. Repeated copies of the same operator are formally distinguished as elements of a graph's operator set and are termed identical operators. Operators that can be interchanged without altering the graph are termed interchangeable operators. This occurs when they are connected to the same set of operators in the same way. When identical operators are also interchangeable, we call them equivalent operators. More generally, if G[O'] denotes the subgraph associated with<sup>3</sup> some subset  $O' \subset O$  of the operators in G, then two disjoint subgraphs<sup>4</sup> can also be classified as identical, interchangeable, or equivalent. For our purposes, these terms will be restricted to connected subgraphs.

**Definition 4.5.** Summand graph. A summand graph<sup>5</sup>  $\Sigma(G) = (G, S, s)$  associates each line l in G with a symbol  $s(l) \in S$  through the label mapping  $s: L \to S$ . Pictorially, this corresponds to labeling each line in G with spin-orbital index, p, q, r, s, etc.  $\Sigma(G)$  translates directly into an algebraic summand according to Def 4.1 and Notation 4.1.

**Definition 4.6.** Degeneracy. The line degeneracy or simply degeneracy of G is the number of permutational symmetries in  $\Sigma(G)$ , a positive integer here denoted  $\mathrm{dg}(G)$ . Formally, this can be defined as follows. If  $S = \{s_1, \ldots, s_n\}$  is the label set of  $\Sigma(G)$  and  $s(l_i) = s_i$  is its label map, we can define a new summand graph  $\Sigma_{\pi}(G) = (G, S, s_{\pi})$  with a permuted label map  $s_{\pi}$  given by  $s_{\pi}(l_i) = s_{\pi(i)}$ , where  $\pi$  is a permutation in  $S_n$ . Then  $\mathrm{dg}(G)$  is the number of  $\Sigma_{\pi}(G)$  that are equal to  $\Sigma(G)$ . Assuming no identical operators, the degeneracy of G is simply given by  $\mathrm{dg}(G) = |L_1|! \cdots |L_h|!$  where  $L_1 \cup \cdots \cup L_h$  partitions its line set into subsets of equivalent lines and  $|L_i|$  denotes the number of elements in  $L_i$ . When the graph does contain identical operators, there is an additional factor of k! for each set  $\{G[O_1], \ldots, G[O_k]\}$  of equivalent subgraphs.

**Axiom 4.1.** Rules of interpretation. The algebraic interpretation of a graph G is obtained from  $\Sigma(G)$  as follows.

- 1. Multiply  $\Sigma(G)$  by  $dg(G)^{-1}$ , the degeneracy factor.
- 2. Sum each index in  $\Sigma(G)$  over its range.

Symbolically, these rules can be stated as follows:  $G = \frac{1}{\deg(G)} \sum_{\text{Einstein}} \Sigma(G)$ .

**Definition 4.7.** Contraction. Formally, a graph contraction is a map  $G \mapsto c(G)$  joining one or more compatible external lines in G. For example, c might replace  $l_1$  and  $l_2$ , which have ends  $(t(l_1), h(l_1)) = (o_1, e)$  and  $(t(l_2), h(l_2)) = (e, o_2)$ , with  $l_{12}$ , which has ends  $(t(l_{12}), h(l_{12})) = (o_1, o_2)$ . Two contractions c and c' of G are equivalent if they are graphically indistinguishable, i.e. c(G) = c'(G). The number of equivalent ways of achieving a given contraction c is called its pattern degeneracy, which is denoted pat(c). The complete set of unique contraction patterns for a graph is here denoted Ctr(G). Combined with Wick's theorem, these concepts lead to the following statement

$$\sum_{\text{Einstein}} \Sigma(G) = \sum_{\text{Einstein}} i\Sigma(G)i + \sum_{c \in \text{Ctr}(G)} \text{pat}(c) \sum_{\text{Einstein}} i\Sigma(c(G))i$$
(4.6)

which will be used to prove a more elegant graphical formulation of Wick's theorem below.

<sup>&</sup>lt;sup>2</sup>In graph theory jargon this is essentially a *directed multigraph*, except that the vertical ordering of operators matters.

 $<sup>^3\</sup>mathrm{See}\ \mathrm{https://en.wikipedia.org/wiki/Induced\_subgraph}$ 

<sup>&</sup>lt;sup>4</sup>That is,  $G[O_1]$  and  $G[O_2]$  with  $O_1 \cap O_2 = \emptyset$ .

<sup>&</sup>lt;sup>5</sup>In graph theory jargon this is an edge-labeled directed multigraph.

**Lemma 4.1.** The pattern degeneracy of a contraction c is  $pat(c) = \frac{dg(G)}{dg(c(G))}$ .

Proof<sup>‡</sup>: Assume G has no identical operators, so that equivalent subgraphs do not contribute to the line degeneracies. If  $L = L_1 \cup \cdots \cup L_m$  partitions G's line set into equivalent lines, let  $L_{i,j} = L_{j,i}$  denote lines in c(G) produced by contracting lines from  $L_i$  and  $L_j$ , and let  $L_{i,0}$  denote lines from  $L_i$  unaffected by c. Then the number of equivalent ways to partition  $L_i$  for contraction is  $\frac{|L_i|!}{|L_{i,0}|!|L_{i,1}|!\cdots|L_{i,m}|!}$ , and the number of ways of forming each  $L_{i,j}$  from a given partition of  $L_i$  and  $L_j$  is  $|L_{i,j}|!$ . Therefore, the total pattern degeneracy of c is given by the following.

$$\operatorname{pat}(c) = \prod_{i=1}^{m} \frac{|L_i|!}{|L_{i,0}|!|L_{i,1}|!\dots|L_{i,m}|!} \cdot \prod_{i>j}^{m} |L_{i,j}|! = \frac{|L_1|!\dots|L_m|!}{\prod_{i=1}^{m} |L_{i,0}|!|L_{i,1}|!\dots|L_{i,i-1}|!} = \frac{\operatorname{dg}(G)}{\operatorname{dg}(c(G))}$$

The final step follows from the fact that, according to our definitions,  $\bigcup_{i=1}^{m} (L_{i,0} \cup L_{i,1} \cup \cdots L_{i,i-1})$  partitions the line set of c(G) into equivalent lines and therefore the denominator equals dg(c(G)).

Remark 4.1. The assumption that G contains no identical operators limits the scope of lemma 4.1 and of the next theorem, which derives from it. We can generalize this result to allow identical operators if we assume that the contraction c does not create any new pairs of equivalent subgraphs.<sup>6</sup> The proof is as follows: Each set  $E = \{O_1, \ldots, O_k\}$  of equivalent subgraph generators<sup>7</sup> for G partitions into  $E_1 \cup \cdots \cup E_n$ , where  $E_i \subset E$  generates a set of equivalent subgraphs in c(G). By our assumption, tallying up these partitions for each E exhausts the sets of equivalent subgraph generators for c(G). Then for each E there will be a factor of |E|! in the line degeneracy of G, a factor of  $|E_1|! \cdots |E_n|!$  in the line degeneracy of C(G), and a factor of C(G) in the pattern degeneracy of the contraction, so that lemma 4.1 still holds.

Theorem 4.1. Wick's theorem for Graphs. 
$$G = :G: + \sum_{c \in Ctr(G)} :c(G): = :G: + \sum_{c \in Ctr(G)}$$

Proof: This follows by expanding G according to axiom 4.1 and then using equation 4.6 and lemma 4.1.

$$G = \frac{1}{\operatorname{dg}(G)} \sum_{\text{Einstein}} \Sigma(G) = \frac{1}{\operatorname{dg}(G)} \sum_{\text{Einstein}} \mathbf{i} \Sigma(G) \mathbf{i} + \sum_{c \in \operatorname{Ctr}(G)} \frac{1}{\operatorname{dg}(c(G))} \sum_{\text{Einstein}} \mathbf{i} \Sigma(c(G)) \mathbf{i} = \mathbf{i} G \mathbf{i} + \sum_{c \in \operatorname{Ctr}(G)} \mathbf{i} c(G) \mathbf{i}$$

**Derivation 4.1.** Phase rule for graphs with contractions. The summand graph of a pair of m-electron operators with a single particle or hole contraction has the following form in the Goldstone representation

where d is a virtual index, k is an occupied index, and we are not using implicit summation. Using the permutational degrees of freedom of the normal-ordered product, we can bring together the contracted pair of single-excitation operators without changing the sign of the expression. The contracted pair of operators can then be eliminated as follows.

$$\mathbf{i} a_{d \bullet}^{p_i} a_{s_i}^{d \bullet} \mathbf{i} = -\mathbf{i} a_{d \bullet s_i}^{d \bullet} \mathbf{i} \mathbf{i} = -(-\eta_d^d) a_{s_i}^{p_i} = a_{s_i}^{p_i}$$

$$\mathbf{i} a_{q_i}^{k \circ} a_{k}^{r_j} \mathbf{i} = -\mathbf{i} a_{k \circ q_i}^{k \circ r_j} \mathbf{i} = -(+\gamma_k^k) a_{q_i}^{r_j} = -a_{q_i}^{r_j}$$

$$(4.8)$$

Applying this rule to each internal line in an open cycle leaves  $(-)^h a_q^p$  where h is the number of hole contractions in the cycle and p and q are indices of the free ends. Applying the same procedure to a loop gives  $(-)^h a_d^{d_{\bullet}} = (-)^h (-\eta_d^d) = (-)^{h+1}$ . Since the lines of any Goldstone graph uniquely partition into Goldstone cycles, these rules provide a quick way to resolve the operator component of any diagram into a normal-ordered product of uncontracted operators times a phase factor.

Corollary 4.1. Phase rule for closed graphs. After eliminating all contracted operator pairs, the sign of a closed graph is  $(-)^{h+l}$ , where h is the total number of holes in the graph and l is the number of loops.

Proof: By definition, every Goldstone cycle in a closed graph is a loop. Let  $h_i$  be the number of holes in the i<sup>th</sup> loop. Then, by derivation 4.1, the fully contracted operator product evaluates to  $\prod_{i=1}^{l} (-)^{h_i+1} = (-)^{\sum_i h_i+l} = (-)^{h+l}$ .

<sup>&</sup>lt;sup>‡</sup>Note that this proof is limited in scope. See rmk 4.1.

<sup>&</sup>lt;sup>6</sup>This encompasses most cases of interest, but it is not completely general, so be cautious about when you apply thm 4.1.

<sup>&</sup>lt;sup>7</sup>By this we mean that  $O_1, \ldots, O_k$  are connected subsets of O such that  $G[O_1], \ldots, G[O_k]$  are equivalent subgraphs.

**Definition 4.8.** Coefficient graph. A coefficient graph is a closed graph in which one of the interaction tensors is an antisymmetrized Kronecker delta,  $\bar{\delta}_{p'_1\cdots p'_m}^{q'_1\cdots q'_m} = \hat{P}_{(q_1/\cdots/q_m)}^{(p_1/\cdots/p_m)} \delta_{p'_1}^{p_1} \cdots \delta_{p'_m}^{p_m} \delta_{q_1}^{q'_1} \cdots \delta_{q_m}^{q'_m}$ . Using the identity

$$\tilde{a}_{q_{1}\dots q_{m}}^{p_{1}\dots p_{m}} = \left(\frac{1}{m!}\right)^{2} \overline{\delta}_{p'_{1}\dots p'_{m}}^{q'_{1}\dots q'_{m}} \tilde{a}_{q'_{1}\dots q'_{m}}^{p'_{1}\dots p'_{m}}$$

$$(4.9)$$

this extends the results derived above to allow for bare excitation operators in addition to m-electron operators. The contraction with  $\tilde{a}_{q_1...q_m}^{p_1...p_m}$  is depicted by capping the lines connected to it,  $\longrightarrow$ , and labelling these creation and annihilation lines with  $p_1, \ldots, p_m$  and  $q_1, \cdots, q_m$ , respectively.<sup>9</sup> To simplify the algebraic interpretation of a coefficient graph (see rmk 4.2), it becomes convenient to treat *coefficient lines* as distinct from other internal lines. A closed graph with no coefficient lines is sometimes called an *energy graph*.

Remark 4.2. The algebraic interpretation of a coefficient graph with the operator in eq 4.9 has the form

$$\bar{\delta}_{p'_1 \cdots p'_m}^{q'_1 \cdots q'_m} T_{q'_1 \cdots q'_m}^{p'_1 \cdots p'_m} = \hat{P}_{(q_1 / \cdots / q_m)}^{(p_1 / \cdots / p_m)} T_{q_1 \cdots q_m}^{p_1 \cdots p_m}$$

$$\tag{4.10}$$

where  $T_{q_1'\cdots q_m'}^{p_1'\cdots p_m'}$  is the product of interaction tensors contracted with  $\overline{\delta}_{p_1'\cdots p_m'}^{q_1'\cdots q_m'}$ , scaled by appropriate sign and degeneracy factors according to axiom 4.1. Suppose  $\{p_1,\ldots,p_m\}$  and  $\{q_1,\cdots,q_m\}$  can be partitioned as  $P_1\cup\cdots\cup P_h$  and  $Q_1\cup\cdots\cup Q_k$  where  $P_i=\{p_{i,1},\ldots,p_{i,|P_i|}\}$  and  $Q_i=\{q_{i,1},\ldots,q_{i,|Q_i|}\}$  label equivalent coefficient lines. Then eq 4.10 simplifies as follows.

$$\hat{P}_{(q_1/\cdots/q_m)}^{(p_1/\cdots/p_m)} T_{q_1\cdots q_m}^{p_1\cdots p_m} = |P_1|!\cdots|P_h|!|Q_1|!\cdots|Q_k|! \,\hat{P}_{(Q_1/\cdots/Q_k)}^{(P_1/\cdots/P_h)} T_{q_1\cdots q_m}^{p_1\cdots p_m} \tag{4.11}$$

Notice that the factorials on the right exactly cancel the contribution of equivalent coefficient lines to the degeneracy factor. This equation defines a simplified rule for interpreting coefficient graphs.

Remark 4.3. Useful results for interpreting graphs. A few results from the preceding discussion are particularly useful for algebraically interpreting a graph according to axiom 4.1. For easy reference, let's summarize them here.

- 1. Each set of k equivalent internal lines<sup>10</sup> or equivalent subgraphs contributes a factor of k! to the degeneracy.
- 2. Each open cycle contributes  $(-)^{h_i}a_q^p$  to the normal-ordered product, where and p and q label the free ends and  $h_i$  is the number of hole contractions in the cycle. A closed cycle (loop) contributes  $(-)^{h_i+1}$ .
- 3. The overall sign of a closed graph is  $(-)^{h+l}$ , where h and l denote the total number of hole lines and loops.
- 4. For each bare excitation operator in a coefficient graph, the coefficient lines contribute an antisymmetrizer  $\hat{P}_{(Q_1/\cdots/Q_k)}^{(P_1/\cdots/P_h)}$  where the  $P_i$ 's and  $Q_i$ 's label subsets of equivalent creation and annihilation lines, respectively.

**Example 4.1.** The one- and two-electron components of  $H_e$  expand into occupied/virtual blocks as follows.

$$h_p^q a_q^p = h_a^b a_b^a + h_a^i a_i^a + h_i^a a_a^i + h_i^j a_j^i$$
(4.12)

$$\frac{1}{4}\overline{g}_{pq}^{rs}a_{rs}^{pq} = \frac{1}{4}\overline{g}_{ab}^{cd}a_{cd}^{ab} + \frac{1}{2}\overline{g}_{ab}^{ci}a_{ci}^{ab} + \frac{1}{2}\overline{g}_{ab}^{bc}a_{bc}^{ai} + \frac{1}{4}\overline{g}_{ab}^{ij}a_{ij}^{ab} + \overline{g}_{ia}^{bj}a_{bj}^{ia} + \frac{1}{4}\overline{g}_{ij}^{ab}a_{ab}^{ij} + \frac{1}{2}\overline{g}_{ia}^{jk}a_{jk}^{ia} + \frac{1}{2}\overline{g}_{ij}^{ka}a_{ka}^{ij} + \frac{1}{4}\overline{g}_{ij}^{kl}a_{kl}^{ij}$$
 (4.13)

which, defining  $\boxtimes \stackrel{\uparrow}{\longrightarrow} \equiv h_p^q a_q^p$  and  $\updownarrow \longrightarrow \updownarrow \equiv \frac{1}{4} \overline{g}_{pq}^{rs} a_{rs}^{pq}$ , can be expressed in terms of Goldstone diagrams as

The degeneracy factors fall into three cases:  $\{l_1, l_2\} \cup \{l_3, l_4\} \implies \operatorname{dg}(G)^{-1} = \frac{1}{2 \cdot 2}; \{l_1, l_2\} \cup \{l_3\} \cup \{l_4\} \implies \operatorname{dg}(G)^{-1} = \frac{1}{2 \cdot 1 \cdot 1};$  and  $\{l_1\} \cup \{l_2\} \cup \{l_3\} \cup \{l_4\} \implies \operatorname{dg}(G)^{-1} = \frac{1}{1 \cdot 1 \cdot 1 \cdot 1}.$  In terms of Hugenholtz diagrams, these are written as follows.

Note that in each case the correct scalar factor is built into the definition of the diagram.

<sup>&</sup>lt;sup>8</sup>The unprimed indices are fixed. This equals the derivative of a generic interaction tensor  $\overline{v}_{p'_1 \cdots p'_m}^{q'_1 \cdots q'_m}$  with respect to  $\overline{v}_{p_1 \cdots p_m}^{q_1 \cdots q_m}$ .

<sup>&</sup>lt;sup>9</sup>If there are multiple bare excitation operators, this notation leads to ambiguities and is best avoided.

<sup>&</sup>lt;sup>10</sup>If rule 4 is used, this excludes coefficient lines. Otherwise, replace rule 4 with the full antisymmetrizer.

**Example 4.2.** The  $\Phi$ -normal Wick expansion of the one- and two-electron components of  $H_e$  are as follows.

$$h_p^q a_q^p = h_p^q \left( \tilde{a}_q^p + \tilde{a}_{q^{\circ}}^{p^{\circ}} \right) = h_p^q \tilde{a}_q^p + h_p^q \gamma_q^p \tag{4.17}$$

$$\frac{1}{4}\overline{g}_{pq}^{rs}a_{rs}^{pq} = \frac{1}{4}\overline{g}_{pq}^{rs}\left(\tilde{a}_{rs}^{pq} + \hat{P}_{(r/s)}^{(p/q)}\tilde{a}_{r^{\circ}s}^{p^{\circ}q} + \hat{P}_{(r/s)}\tilde{a}_{r^{\circ}s^{\circ\circ}}^{p^{\circ}q^{\circ\circ}}\right) = \frac{1}{4}\overline{g}_{pq}^{rs}\tilde{a}_{rs}^{pq} + \overline{g}_{pq}^{rs}\gamma_{r}^{p}\tilde{a}_{s}^{q} + \frac{1}{2}\overline{g}_{pq}^{rs}\gamma_{r}^{p}\gamma_{s}^{q}$$

$$(4.18)$$

In terms of Goldstone diagrams and Hugenholtz diagrams, these equations are written as follows.

$$\updownarrow \qquad \qquad \updownarrow \qquad \qquad (4.20)$$

$$\updownarrow \qquad \qquad \updownarrow \qquad \qquad \updownarrow \qquad \qquad (4.20)$$

$$\Diamond = \Diamond + \bigcirc \qquad \qquad \qquad (4.21)$$

Again each degeneracy factor is exactly equal to the scalar factor in front of the corresponding algebraic term, in accord with Wick's theorem for graphs. Using these results, the  $\Phi$ -normal Wick expansion of  $H_e$  in Goldstone representation is

where  $\otimes$ — $\stackrel{\diamond}{\phi}$  is the Fock operator,  $f_p^q \tilde{a}_q^p = h_p^q \tilde{a}_q^p + \overline{g}_{pr}^{qs} \gamma_s^r \tilde{a}_q^p$ 

**Notation 4.2.** The bra and ket of the reference state  $\Phi$  are graphically depicted by thick double lines  $\longrightarrow$  above and below the graph, respectively. Open graphs with a ket at the bottom are sometimes classified as wavefunction graphs.

Remark 4.4. The singles, doubles, triples, etc. configuration interaction (CI) operators will here be denoted as follows.

$$C_1 \equiv c_a^i \tilde{a}_i^a \equiv \bigvee \qquad C_2 \equiv (\frac{1}{2!})^2 c_{ab}^{ij} \tilde{a}_{ij}^{ab} \equiv \bigvee \qquad C_3 \equiv (\frac{1}{3!})^2 c_{abc}^{ijk} \tilde{a}_{ijk}^{abc} \equiv \bigvee \qquad \cdots$$
 (4.23)

CI theory uses these operators to define a linear parametrization of the wavefunction,  $\Psi_{\text{CI}} = (1 + C_1 + \dots + C_n)\Phi^{1}$ . Coupled-cluster (CC) theory employs a similar series of operators  $T_k \equiv (\frac{1}{k!})^2 t_{a_1 \dots a_k}^{i_1 \dots i_k} \tilde{a}_{i_1 \dots i_k}^{a_1 \dots a_k}$  to define an exponential parametrization  $\Psi_{\rm CC} = \exp(T_1 + \cdots + T_n)\Phi$ .

Example 4.3. Derivation of CIS in diagram notation. The CI singles transition energy to an excited state is  $\omega \equiv E - E_0 = \langle \Phi | C_1^{\dagger} H_c C_1 | \Phi \rangle$ . Using Wick's theorem for graphs, we can evaluate this expectation value as

$$\omega = \boxed{\left( \bigotimes_{j}^{b} + \overset{\dagger}{\downarrow} \bigotimes_{j}^{b} \right)} = \bigotimes_{j}^{a} + \bigotimes_{j}^{b} + \bigotimes_{j}^{a} c_{j}^{b} = c_{i}^{a} (f_{a}^{b} \delta_{j}^{i} - f_{j}^{i} \delta_{a}^{b} + \overline{g}_{ja}^{bi}) c_{b}^{j}$$

where the expression in parentheses on the right defines the CIS matrix,  $\langle \Phi_i^a | H_c | \Phi_i^b \rangle$ , which we could have also evaluated separately as a coefficient graph. The main consideration in evaluating such expectation values is to identify which terms in the quasiparticle expansion of  $H_c$  can form complete contractions with the CI operators in the bra and the ket.

Example 4.4. Derivation of the ground-state CI energy. Projecting the CI Schrödinger equation  $^{12}$  by  $\Phi$ , the exact ground-state correlation energy can be expressed in CI as  $E_c = \langle \Phi | H_c (1 + C_1 + \cdots + C_n) | \Phi \rangle$ . Using WTG, we find

$$E_{c} = \underbrace{\begin{pmatrix} \begin{pmatrix} \begin{pmatrix} -\frac{1}{2} + \frac{1}{2} \end{pmatrix} \end{pmatrix} \begin{pmatrix} \begin{pmatrix} -\frac{1}{2} + \frac{1}{2} \end{pmatrix} \end{pmatrix} \begin{pmatrix} -\frac{1}{2} \end{pmatrix} \begin{pmatrix}$$

where the first term evaluates to zero whenever Brillouin's theorem applies. This shows that the CI energy depends on triples, quadruples, etc. only indirectly, through their influence on  $C_1$  and  $C_2$ .

<sup>&</sup>lt;sup>11</sup>This expansion assumes "intermediate normalization", in which we normalize the wavefunction to satisfy  $\langle \Phi | \Psi \rangle = 1$ . In standard normalization, the CI wavefunction is given by  $\Psi_{\text{CI}} = (c_0 + C_1 + \dots + C_n)\Phi$ , where  $c_0$  is the scalar weight of the reference function.

**Example 4.5.** Derivation of the CID coefficient equation. Projecting the CID Schrödinger equation by  $\Phi_{ij}^{ab}$  leads to the CID coefficient equation,  $\langle \Phi_{ij}^{ab} | H_c(1+C_2) | \Phi \rangle = E_c c_{ab}^{ij}$ . The left-hand side can be evaluated as follows.<sup>13</sup>

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Note that the equivalent coefficient lines in these diagrams don't contribute to the degeneracy factors here, since these have been canceled using item 4 in rmk 4.3. The minus sign in the third term comes from the fact that its diagram has three holes and two loops,  $(-1)^{3+2} = -1$ .

Example 4.6. Derivation of the CCD amplitude equation. Projecting the CCD Schrödinger equation by  $\Phi_{ij}^{ab}$  leads to the CCD amplitude equation,  $\langle \Phi_{ij}^{ab} | H_c \exp(T_2) | \Phi \rangle = E_c t_{ab}^{ij}$ . Note that terms cubic and higher in  $T_2$  cannot form complete contractions with  $\tilde{a}_{ab}^{ij} H_c$ , reducing the left-hand side to  $\langle \Phi_{ij}^{ab} | H_c (1 + T_2 + \frac{1}{2}T_2^2) | \Phi \rangle$ . If we exchange CI coefficients for CC amplitudes the linear terms are given by equation 4.24, so only the quadratic term remains to be evaluated.

$$\tfrac{1}{2} \langle \Phi^{ab}_{ij} | H_{\text{c}} \, T_2^2 | \Phi \rangle = (\tfrac{1}{2})^2 \, \overline{g}^{cd}_{kl} t^{kl}_{ab} t^{ij}_{cd} + \tfrac{1}{2} \hat{P}^{(a/b)}_{(i/j)} \overline{g}^{cd}_{kl} t^{ik}_{ac} t^{lj}_{db} - \tfrac{1}{2} \hat{P}_{(a/b)} \, \overline{g}^{cd}_{kl} t^{kl}_{ca} t^{ij}_{db} - \tfrac{1}{2} \hat{P}^{(i/j)} \overline{g}^{cd}_{kl} t^{ki}_{cd} t^{lj}_{ab} + E_{\text{c}} \, t^{ij}_{ab} + E_{\text{c}} \, t^{ij}_{$$

In the last term, we have substituted in the CCD energy expression, which is the same graph as the CID energy expression.

$$E_{\rm c} = \underbrace{\begin{pmatrix} \bigotimes \downarrow \downarrow \downarrow \downarrow \downarrow \downarrow \\ \downarrow \downarrow \downarrow \downarrow \downarrow \end{pmatrix}}_{=} = \underbrace{\begin{pmatrix} \frac{1}{2} \end{pmatrix}^2 \overline{g}_{ij}^{ab} t_{ab}^{ij}}_{=b}$$

$$(4.25)$$

Note that the term  $E_c t_{ab}^{ij}$  actually cancels with the right-hand side of the projected Schrödinger equation. Since this is the only unlinked graph in the expansion, we can write the amplitude equation as

$$\langle \Phi_{ij}^{ab} | H_{c} \exp(T_2) | \Phi \rangle_{L} = 0 \tag{4.26}$$

where the subscript indicates that only linked graphs are included. This cancellation of unlinked terms is a general feature of coupled-cluster theory. Finally, note that when Brillouin's theorem holds the second and third terms in eq 4.24 become

$$\hat{P}_{(a/b)}f_a^c t_{cb}^{ij} - \hat{P}^{(i/j)}f_k^i t_{ab}^{kj} = -(\epsilon_i + \epsilon_j - \epsilon_a - \epsilon_b)t_{ab}^{ij}$$

where  $\{\epsilon_p = f_p^p\}$  are canonical Hartree-Fock orbital energies. Subtracting this contribution from both sides leads to

$$t_{ab}^{ij} = (\mathcal{E}_{ab}^{ij})^{-1} \langle \Phi_{ij}^{ab} | (H_c - f_p^p \tilde{a}_p^p) \exp(T_2) | \Phi \rangle_{\mathcal{L}} \qquad \qquad \mathcal{E}_{ab}^{ij} \equiv \epsilon_i + \epsilon_j - \epsilon_a - \epsilon_b \qquad (4.27)$$

which is the standard form for the  $T_2$  amplitude equation.

**Definition 4.9.** Resolvent line. Complete contractions through a resolvent line: are defined as follows.

$$\mathbf{i} a^{i_1} \cdots a^{i_k} \overline{a_{a_1} \cdots a_{a_k}} a^{b_1} \cdots a^{b_k} a_{j_1} \cdots a_{j_k} \mathbf{i} \equiv \frac{\mathbf{i} a^{i_1} \cdots a^{i_k} \overline{a_{a_1} \cdots a_{a_k}} a^{b_1} \cdots a^{b_k} a_{j_1} \cdots a_{j_k} \mathbf{i}}{\mathcal{E}^{i_1 \cdots i_k}_{a_1 \cdots a_k}}$$
 
$$\mathcal{E}^{i_1 \cdots i_k}_{a_1 \cdots a_k} \equiv \sum_{r=1}^k \epsilon_{i_r} - \sum_{r=1}^k \epsilon_{a_r} a^{b_1} \cdots a^{b_r} a^{b_r}$$

This can be denoted in a graph by drawing a horizontal dotted line through several contraction lines.

<sup>&</sup>lt;sup>12</sup>By this we mean  $H_c\Psi_{CI} = E_c\Psi_{CI}$ , where  $E_c = E - E_0$  is the correlation energy.

<sup>&</sup>lt;sup>13</sup>Note that we are using Einstein summation here, so c, d, k, and l are summation indices.

**Example 4.7.** Using the notion of a resolvent line, equation 4.27 can be written in diagram notation as follows.

$$\begin{array}{c} \begin{array}{c} \\ \\ \\ \end{array} \end{array} \begin{array}{c} \\ \\ \end{array}$$

This equation is typically solved in an iterative procedure that updates the amplitudes by substituting the previous iteration's amplitudes into the right-hand side, starting with  ${}^{[0]}t^{ij}_{ab}=0$  as an initial guess.

Remark 4.5. The  $CEPA_0$  and MP2 amplitude equations. If the  $T_2$  amplitudes are small, the CCD exponential,  $\exp(T_2) = 1 + T_2 + \frac{1}{2}T_2^2 + \ldots$ , can be approximated by the linear terms or even just the constant term, leading to the coupled electron-pair approximation zero ( $CEPA_0$ ) and second-order  $M\emptyset$ ller-Plesset perturbation theory (MP2) methods. Assuming Brillouin's theorem and writing V for the electron repulsion operator,  $\frac{1}{4}\overline{g}_{pq}^{rs}\tilde{a}_{rs}^{pq}$ , these amplitude equations are

where the MP2 amplitude is usually denoted  ${}^{(1)}t^{ij}_{ab}$  because it is correct to first order in perturbation theory. Note that eq 4.29 specifies the MP2 amplitudes directly,  ${}^{(1)}t^{ij}_{ab}=(\mathcal{E}^{ij}_{ab})^{-1}\overline{g}^{ij}_{ab}$ , and requires no iterative solution. Substituting these amplitudes into equation 4.25 gives the MP2 energy, which has contributions up to second order in perturbation theory.

$$E_{c}^{(2)} = \underbrace{\downarrow}_{(1)} \underbrace{\downarrow}_{(1)} = \underbrace{\downarrow}_{(1)} \underbrace{\downarrow}_{(1)} = \underbrace{\downarrow}_{ijab} \underbrace{\frac{\overline{g}_{ab}^{ij} \overline{g}_{ab}^{ab}}{\mathcal{E}_{ab}^{ij}}}_{(4.31)}$$