On the Independence of Fundamental Decompositions of Power-Law Kinetic Systems

Bryan S. Hernandez\textsuperscript{1,2,*}

\textsuperscript{1}Institute of Mathematical Sciences and Physics, University of the Philippines, Los Baños, Laguna 4031, Philippines
\textsuperscript{2}Institute of Mathematics, University of the Philippines Diliman, Quezon City 1101, Philippines
*Corresponding author: bryan.hernandez@upd.edu.ph

January 16, 2020

Abstract

The fundamental decomposition of a chemical reaction network (CRN) is induced by partitioning the reaction set into “fundamental classes”. It was the basis of the Higher Deficiency Algorithm for mass action systems of Ji and Feinberg, and the Multistationarity Algorithm for power-law kinetic systems of Hernandez et al. In this paper, we provide important properties of the independence (i.e., the network’s stoichiometric subspace is the direct sum of the subnetworks’ stoichiometric subspaces) and the incidence-independence (i.e., the image of the network’s incidence map is the direct sum of the incidence maps’ images of the subnetworks) of these decompositions. Feinberg established the essential relationship between independent decompositions and the set of positive equilibria of a network, which we call the Feinberg Decomposition Theorem (FDT). In addition, Fariñas et al. recently documented its version for incidence-independence. Fundamental decomposition divides the network into subnetworks of deficiency either 0 or 1. Hence, available results for lower deficiency networks, such as the Deficiency Zero Theorem (DZT), can be used. These justify the study of independent fundamental decompositions. A MATLAB program which (i) computes the subnetworks of a CRN under the fundamental decomposition and (ii) determines whether the decomposition is independent and incidence-independent is also created. Finally, we consider the CRN of the generalization of a subnetwork of Schmitz’s carbon cycle model by Hernandez et al. endowed with mass action kinetics, and we solve the problem of its multistationarity using (1) the program, (2) the DZT, and (3) the FDT.
1 Introduction

The fundamental decomposition (or “$F$-decomposition”) of a chemical reaction network (CRN) is the set of subnetworks generated by the partition of its set of reactions into “fundamental classes”. This was introduced by Ji and Feinberg [12] in 2011 as the basis of their Higher Deficiency Algorithm (HDA) for mass action systems. In 2020, Hernandez et al. [10] extended the HDA to power-law kinetic systems with reactant-determined interactions (PL-RDK). These are reactions branching from the same reactant complex having identical kinetic order vectors. By combining this extension with a method that transforms a power-law kinetic system with non-reactant-determined interactions (PL-NDK) to a dynamically equivalent PL-RDK system, the Multistationarity Algorithm (MSA) was established and used to determine multistationarity (i.e., the system admits at least two equilibria) of any power-law kinetic system within a stoichiometric class.

This paper explores the properties of the $F$-decomposition, in particular, its independence (i.e., the network’s stoichiometric subspace is the direct sum of the subnetworks’ stoichiometric subspaces) and its incidence-independence (i.e., the image of the network’s incidence map is the direct sum of the incidence maps’ images of the subnetworks). In our previous work [11], we mention that for independent fundamental decompositions, the transformation that converts a PL-NDK system to a PL-RDK system is not necessary.

M. Feinberg established the essential relationship between independent decompositions and the set of positive equilibria of a network in 1987 [5]. A corresponding relationship between incidence-independent, weakly reversible decompositions and complex-balanced equilibria of a weakly reversible network was recently documented by Fariñas et al. [4]. These also motivate the study of independent decompositions. Moreover, fundamental decompositions decompose a CRN into subnetworks where each of the subnetworks has deficiency either 0 or 1. Thus, the well-known Deficiency Zero and Deficiency One Theorems, and other lower deficiency theorems in literature, maybe used.

The paper is organized as follows: Section 2 provides the fundamentals of chemical reaction networks (CRNs) and chemical kinetic systems. It also presents relevant results from the decomposition theory of CRNs. Section 3 collects important properties of the fundamental and other related decompositions. In Section 4, we provide results between the independence and the incidence-independence of the fundamental decomposition and the types of the networks. In addition, the solution of the multistationarity of the CRN
of the generalization of a subnetwork of Schmitz’s carbon cycle model by Hernandez et al. endowed with mass action kinetics is also given. In Section 5, we establish relationships between the fundamental decomposition of a kinetic system and its transform under the CF-RM. Conclusions and an outlook constitute Section 6. Tables of acronyms and frequently used symbols are provided in Appendix A. Finally, Appendix B provides a program that outputs the subnetworks under the fundamental decomposition, and determines whether the decomposition is independent or incidence-independent.

2 Fundamentals of Chemical Reaction Networks and Kinetic Systems

In this section, we recall some fundamental notions about chemical reaction networks and chemical kinetic systems [1,6]. We also present important preliminaries on the decomposition theory which was introduced by Feinberg in [5].

2.1 Fundamentals of Chemical Reaction Networks

Definition 2.1. A chemical reaction network (CRN) \( \mathcal{N} \) is a triple \((\mathcal{S}, \mathcal{C}, \mathcal{R})\) of nonempty finite sets where \( \mathcal{S} \), \( \mathcal{C} \), and \( \mathcal{R} \) are the sets of \( m \) species, \( n \) complexes, and \( r \) reactions, respectively, such that \((C_i, C_j) \notin \mathcal{R}\) for each \( C_i \in \mathcal{C} \); and for each \( C_i \in \mathcal{C} \), there exists \( C_j \in \mathcal{C} \) such that \((C_i, C_j) \in \mathcal{R}\) or \((C_j, C_i) \in \mathcal{R}\).

Definition 2.2. The molecularity matrix, denoted by \( Y \), is an \( m \times n \) matrix such that \( Y_{ij} \) is the stoichiometric coefficient of species \( X_i \) in complex \( C_j \). The incidence matrix, denoted by \( I_a \), is an \( n \times r \) matrix such that

\[
(I_a)_{ij} = \begin{cases} 
-1 & \text{if } C_i \text{ is in the reactant complex of reaction } R_j, \\
1 & \text{if } C_i \text{ is in the product complex of reaction } R_j, \\
0 & \text{otherwise.}
\end{cases}
\]

The stoichiometric matrix, denoted by \( N \), is the \( m \times r \) matrix given by \( N = Y I_a \).

Let \( \mathcal{I} = \mathcal{I}, \mathcal{C} \) or \( \mathcal{R} \). We denote the standard basis for \( \mathbb{R}^\mathcal{I} \) by \( \{ \omega_i \in \mathbb{R}^\mathcal{I} \mid i \in \mathcal{I} \} \).

Definition 2.3. Let \( \mathcal{N} = (\mathcal{I}, \mathcal{C}, \mathcal{R}) \) be a CRN. The incidence map \( I_a : \mathbb{R}^\mathcal{I} \to \mathbb{R}^\mathcal{C} \) is the linear map such that for each reaction \( r : C_i \to C_j \in \mathcal{R} \), the basis vector \( \omega_r \) to the vector \( \omega_{C_j} - \omega_{C_i} \in \mathcal{C} \).

Definition 2.4. The reaction vectors for a given reaction network \((\mathcal{I}, \mathcal{C}, \mathcal{R})\) are the elements of the set \( \{ C_j - C_i \in \mathbb{R}^\mathcal{I} \mid (C_i, C_j) \in \mathcal{R} \} \).
Definition 2.5. The stoichiometric subspace of a reaction network \((\mathcal{S}, \mathcal{C}, \mathcal{R})\), denoted by \(S\), is the linear subspace of \(\mathbb{R}^\mathcal{S}\) given by 
\[
S = \text{span} \{ C_j - C_i \in \mathbb{R}^\mathcal{S} | (C_i, C_j) \in \mathcal{R} \}
\].
The rank of the network, denoted by \(s\), is given by \(s = \dim S\). The set \((x + S) \cap \mathbb{R}_{\geq 0}^\mathcal{S}\) is said to be a stoichiometric compatibility class of \(x \in \mathbb{R}_{\geq 0}^\mathcal{S}\).

Definition 2.6. Two vectors \(x, x^* \in \mathbb{R}^\mathcal{S}\) are stoichiometrically compatible if \(x - x^*\) is an element of the stoichiometric subspace \(S\).

We can view complexes as vertices and reactions as edges. With this, CRNs can be seen as graphs. At this point, if we are talking about geometric properties, vertices are complexes and edges are reactions. If there is a path between two vertices \(C_i\) and \(C_j\), then they are said to be connected. If there is a directed path from vertex \(C_i\) to vertex \(C_j\) and vice versa, then they are said to be strongly connected. If any two vertices of a subgraph are (strongly) connected, then the subgraph is said to be a (strongly) connected component. The (strong) connected components are precisely the (strong) linkage classes of a CRN. The maximal strongly connected subgraphs where there are no edges from a complex in the subgraph to a complex outside the subgraph is said to be the terminal strong linkage classes. We denote the number of linkage classes and the number of strong linkage classes by \(l\) and \(sl\), respectively. A CRN is said to be weakly reversible if \(sl = l\).

Definition 2.7. For a CRN, the deficiency is given by \(\delta = n - l - s\) where \(n\) is the number of complexes, \(l\) is the number of linkage classes, and \(s\) is the dimension of the stoichiometric subspace \(S\).

2.2 Fundamentals of Chemical Kinetic Systems

Definition 2.8. A kinetics \(K\) for a reaction network \((\mathcal{S}, \mathcal{C}, \mathcal{R})\) is an assignment to each reaction \(r : y \to y' \in \mathcal{R}\) of a rate function \(K_r : \Omega_K \to \mathbb{R}_{\geq 0}\) such that \(\mathbb{R}_{\geq 0} \subseteq \Omega_K \subseteq \mathbb{R}^\mathcal{S}\), \(c \wedge d \in \Omega_K\) if \(c, d \in \Omega_K\), and \(K_r(c) \geq 0\) for each \(c \in \Omega_K\). Furthermore, it satisfies the positivity property: \(\text{supp } y \subseteq \text{supp } c\) if and only if \(K_r(c) > 0\). The system \((\mathcal{S}, \mathcal{C}, \mathcal{R}, K)\) is called a chemical kinetic system.

Definition 2.9. The species formation rate function (SFRF) of a chemical kinetic system is given by 
\[
f(x) = NK(x) = \sum_{C_i \to C_j \in \mathcal{R}} K_{C_i \to C_j}(x) (C_j - C_i).
\]
The ordinary differential equation (ODE) or dynamical system of a chemical kinetics system is \( \frac{dx}{dt} = f(x) \). An equilibrium or steady state is a zero of \( f \).

**Definition 2.10.** The set of positive equilibria of a chemical kinetic system \((\mathcal{S}, \mathcal{C}, \mathcal{R}, K)\) is given by \( E_+ (\mathcal{S}, \mathcal{C}, \mathcal{R}, K) = \{ x \in \mathbb{R}_{>0}^S | f(x) = 0 \} \).

A CRN is said to admit multiple equilibria if there exist positive rate constants such that the ODE system admits more than one stoichiometrically compatible equilibria.

**Definition 2.11.** A kinetics \( K \) is complex factorizable if, for \( K(x) = k I_K(x) \), the interaction map \( I_K : \mathbb{R}^S \rightarrow \mathbb{R}^R \) factorizes via the space of complexes \( \mathbb{R}^C : I_K = I_k \circ \psi_K \) with \( \psi_K : \mathbb{R}^S \rightarrow \mathbb{R}^C \) as factor map and \( I_k = \text{diag}(k) \circ \rho' \) with \( \rho' : \mathbb{R}^C \rightarrow \mathbb{R}^R \) assigning the value at a reactant complex to all its reactions.

**Definition 2.12.** A kinetics \( K \) is a power-law kinetics (PLK) if \( K_i(x) = k_i x^{F_i} \) for \( i = 1, ..., r \) where \( k_i \in \mathbb{R}_{>0} \) and \( F_{ij} \in \mathbb{R} \). The power-law kinetics is defined by an \( r \times m \) matrix \( F \), called the kinetic order matrix and a vector \( k \in \mathbb{R}^R \), called the rate vector.

If the kinetic order matrix is the transpose of the molecularity matrix, then the system becomes the well-known mass action kinetics (MAK).

**Definition 2.13.** A PLK system has reactant-determined kinetics (of type PL-RDK) if for any two reactions \( i, j \) with identical reactant complexes, the corresponding rows of kinetic orders in \( F \) are identical, i.e., \( f_{ik} = f_{jk} \) for \( k = 1, 2, ..., m \). A PLK system has non-reactant-determined kinetics (of type PL-NDK) if there exist two reactions with the same reactant complexes whose corresponding rows in \( F \) are not identical.

We now state the Deficiency Zero and Deficiency One Theorems by Feinberg [5–7].

**Theorem 2.14.** (Deficiency Zero Theorem) For any CRN of deficiency zero, the following statements hold:

i. If the network is not weakly reversible, then for arbitrary kinetics, the differential equations for the corresponding reaction system cannot admit an equilibrium.

ii. If the network is not weakly reversible, then for arbitrary kinetics, the differential equations for the corresponding reaction system cannot admit a cyclic composition trajectory containing a positive composition.
iii. If the network is weakly reversible, then for any mass action kinetics (but regardless of the positive values the rate constants take), the resulting differential equations have the following properties:
There exists within each positive stoichiometric compatibility class precisely one equilibrium; that equilibrium is asymptotically stable; there is no nontrivial cyclic composition trajectory along which all species concentrations are positive.

**Theorem 2.15. (Deficiency One Theorem)** Consider a mass action system for which the underlying reaction network has $l$ linkage classes. Let $\delta$ be the deficiency of the network and let $\delta_\theta$ be the deficiency of the $\theta$th linkage class. Suppose that the following conditions hold:

i. $\delta_\theta \leq 1$ for each linkage class and

ii. the sum of the deficiencies of all the individual linkage classes is the deficiency of the whole network.

If the network is weakly reversible, then for any mass action kinetics, the differential equations for the system admit precisely one equilibrium in each positive stoichiometric compatibility class.

### 2.3 Review of Decomposition Theory

We now recall some definitions and earlier results from the decomposition theory of chemical reaction networks.

**Definition 2.16.** A decomposition of $\mathcal{N}$ is a set of subnetworks $\{\mathcal{N}_1, \mathcal{N}_2, ..., \mathcal{N}_k\}$ of $\mathcal{N}$ induced by a partition $\{\mathcal{R}_1, \mathcal{R}_2, ..., \mathcal{R}_k\}$ of its reaction set $\mathcal{R}$.

We denote a decomposition with $\mathcal{N} = \mathcal{N}_1 \cup \mathcal{N}_2 \cup ... \cup \mathcal{N}_k$ since $\mathcal{N}$ is a union of the subnetworks in the sense of Section 9. It also follows immediately that, for the corresponding stoichiometric subspaces, $S = S_1 + S_2 + ... + S_k$. It is also useful to consider refinements and coarsenings of decompositions.

**Definition 2.17.** A network decomposition $\mathcal{N} = \mathcal{N}_1 \cup \mathcal{N}_2 \cup ... \cup \mathcal{N}_k$ is a refinement of $\mathcal{N} = \mathcal{N'}_1 \cup \mathcal{N'}_2 \cup ... \cup \mathcal{N'}_{k'}$ (and the latter a coarsening of the former) if it is induced by a refinement $\{\mathcal{R}_1, \mathcal{R}_2, ..., \mathcal{R}_k\}$ of $\{\mathcal{R'}_1 \cup \mathcal{R'}_2 \cup ... \cup \mathcal{R'}_{k'}\}$, i.e., each $\mathcal{R}_i$ is contained in an $\mathcal{R'}_j$. 
In [5], Feinberg introduced the important concept of independent decomposition.

**Definition 2.18.** A network decomposition $\mathcal{N} = \mathcal{N}_1 \cup \mathcal{N}_2 \cup ... \cup \mathcal{N}_k$ is **independent** if its stoichiometric subspace is a direct sum of the subnetwork stoichiometric subspaces.

For any decomposition, it also holds that $\text{Im } I_a = \text{Im } I_{a,1} + ... + \text{Im } I_{a,k}$, where $\text{Im } I_{a,i} = I_a(\mathbb{R}^{\mathcal{N}_i})$.

**Definition 2.19.** A network decomposition is **incidence-independent** if $\text{Im } I_a$ is a direct sum of the $\text{Im } I_{a,i}$. It is **bi-independent** if it is both independent and incidence-independent.

An equivalent formulation of showing incidence-independent is to satisfy $n - l = \sum (n_i - l_i)$, where $n_i$ is the number of complexes and $l_i$ is the number of linkage classes, in each subnetwork $i$.

Feinberg established the following relation between an independent decomposition and the set of positive equilibria of a kinetics on the network.

**Theorem 2.20.** (Feinberg Decomposition Theorem [5]) Let $P(\mathcal{R}) = \{\mathcal{R}_1, \mathcal{R}_2, ..., \mathcal{R}_k\}$ be a partition of a CRN $\mathcal{N}$ and let $K$ be a kinetics on $\mathcal{N}$. If $\mathcal{N} = \mathcal{N}_1 \cup \mathcal{N}_2 \cup ... \cup \mathcal{N}_k$ is the network decomposition of $P(\mathcal{R})$ and $E_+ (\mathcal{N}, K_i) = \{ x \in \mathbb{R}_{>0}^{\mathcal{N}} | \mathcal{N}_i K_i(x) = 0 \}$ then

$$E_+ (\mathcal{N}_1, K_1) \cap E_+ (\mathcal{N}_2, K_2) \cap ... \cap E_+ (\mathcal{N}_k, K_k) \subseteq E_+ (\mathcal{N}, K).$$

If the network decomposition is independent, then equality holds.

The following theorem is the analogue of Feinberg’s result for incidence-independent decompositions and complex-balanced equilibria [4].

**Theorem 2.21.** Let $\mathcal{N}$ be a network, $K$ any kinetics and $\mathcal{N} = \mathcal{N}_1 \cup \mathcal{N}_2 \cup ... \cup \mathcal{N}_k$ an incidence-independent decomposition of weakly reversible subnetworks. Then $\mathcal{N}$ is weakly reversible and

i. $Z_+ (\mathcal{N}, K) = \bigcap Z_+ (\mathcal{N}_i, K)$ for each subnetwork $\mathcal{N}_i$.

ii. If $Z_+ (\mathcal{N}, K) \neq \emptyset$ then $Z_+ (\mathcal{N}_i, K) \neq \emptyset$ for each subnetwork $\mathcal{N}_i$.

iii. If the decomposition is a C-decomposition and $K$ a complex factorizable kinetics then $Z_+ (\mathcal{N}_i, K) \neq \emptyset$ for each subnetwork $\mathcal{N}_i$ implies that $Z_+ (\mathcal{N}, K) \neq \emptyset$. 
3 The $\mathcal{O}$-, $\mathcal{P}$-, and $\mathcal{F}$-decompositions of CRNs

We review the concepts and properties underlying HDA and its extension to PL-RDK systems in the context of decomposition theory \[10,12\].

**Definition 3.1.** A subset $\mathcal{O}$ of $\mathcal{R}$ is said to be an **orientation** if for every reaction $y \rightarrow y' \in \mathcal{R}$, either $y \rightarrow y' \in \mathcal{R}$ or $y' \rightarrow y \in \mathcal{R}$, but not both.

For an orientation $\mathcal{O}$, we define a linear map $L_\mathcal{O} : \mathbb{R}^\mathcal{O} \rightarrow S$ such that

$$L_\mathcal{O}(\alpha) = \sum_{y \rightarrow y' \in \mathcal{O}} \alpha_{y \rightarrow y'} (y' - y).$$

Each orientation $\mathcal{O}$ defines a partition of $\mathcal{N}$ into $\mathcal{O}$ and its complement $\mathcal{O}'$, which generates the following decomposition:

**Definition 3.2.** For an orientation $\mathcal{O}$ on $\mathcal{N}$, the **$\mathcal{O}$-decomposition** of $\mathcal{N}$ consists of the subnetworks $\mathcal{N}_\mathcal{O}$ and $\mathcal{N}_{\mathcal{O}'}$, i.e., $\mathcal{N} = \mathcal{N}_\mathcal{O} \cup \mathcal{N}_{\mathcal{O}'}$.

We now review the important concept of “equivalence classes” from \[12\]. Let $\{v^l\}_{l=1}^d$ be a basis for $\text{Ker} L_\mathcal{O}$. If for $y \rightarrow y' \in \mathcal{O}$, $v^l_{y \rightarrow y'} = 0$ for all $1 \leq l \leq d$ then the reaction $y \rightarrow y'$ belongs to the zeroth equivalence class $P_0$. For $y \rightarrow y', \overline{y} \rightarrow \overline{y}' \in \mathcal{O}' \setminus P_0$, if there exists $\alpha \neq 0$ such that $v^l_{y \rightarrow y'} = \alpha v^l_{\overline{y} \rightarrow \overline{y}'}$ for all $1 \leq l \leq d$, then the two reactions are in the same equivalence class denoted by $P_i$, $i \neq 0$.

The central concept of “fundamental classes” is actually the basis of the Higher Deficiency Algorithm of Ji and Feinberg. The reactions $y \rightarrow y'$ and $\overline{y} \rightarrow \overline{y}'$ in $\mathcal{R}$ belong to the same **fundamental class** if at least one of the following is satisfied \[12\].

i. $y \rightarrow y'$ and $\overline{y} \rightarrow \overline{y}'$ are the same reaction.

ii. $y \rightarrow y'$ and $\overline{y} \rightarrow \overline{y}'$ are reversible pair.

iii. Either $y \rightarrow y'$ or $y' \rightarrow y$, and either $\overline{y} \rightarrow \overline{y}'$ or $\overline{y}' \rightarrow \overline{y}$ are in the same equivalence class on $\mathcal{O}$.

It is worth mentioning that he orientation $\mathcal{O}$ is partitioned into equivalence classes while the reaction set $\mathcal{R}$ is partitioned into fundamental classes.

**Definition 3.3.** The **$\mathcal{F}$-decomposition** of $\mathcal{N}$ is the decomposition generated by the partition of $\mathcal{R}$ into fundamental classes.
Theorem 3.4. [11] Let $\mathcal{N}_\mathcal{O}$ be the subnetwork of $\mathcal{N}$ defined by the orientation $\mathcal{O}$ being a subset of $\mathcal{R}$. Then the following holds:

i. The $\mathcal{P}$-decomposition of $\mathcal{N}_\mathcal{O}$ is independent if and only if the $\mathcal{F}$-decomposition of $\mathcal{N}$ is independent.

ii. The $\mathcal{P}$-decomposition of $\mathcal{N}_\mathcal{O}$ is incidence-independent if and only if the $\mathcal{F}$-decomposition of $\mathcal{N}$ is incidence-independent.

iii. The $\mathcal{P}$-decomposition of $\mathcal{N}_\mathcal{O}$ is bi-independent if and only if the $\mathcal{F}$-decomposition of $\mathcal{N}$ is bi-independent.

We denote the zeroth equivalence class as $P_0$, the nontrivial equivalence classes as $P_1, P_2, ..., P_{w_1}$ and the trivial equivalence classes as $P_{w_1+1}, P_{w_1+2}, ..., P_{w_1+w_2}$. If the zeroth equivalence class is nonempty, then there are $w_1 + w_2 + 1 = w + 1$ equivalence classes where $w = w_1 + w_2$.

We then get the sets $(C_i \setminus P_i)$ for $i = 0, 1, 2, ..., w$. Let $t$ be the number of nonempty sets excluding $P_0$. If $P_0$ is nonempty, we define the $\tilde{\mathcal{F}}$-decomposition in the following manner:

$$\left( \bigcup_{i=0}^{w} P_i \right) \cup \left( \bigcup_{i=0}^{t} (C_i \setminus P_i) \right).$$

We now establish our basic new results for the relationship of $\tilde{\mathcal{P}}$- and $\mathcal{F}$-decompositions.

Proposition 3.5. The $\tilde{\mathcal{P}}$-decomposition is a refinement of the $\mathcal{F}$-decomposition.

Proof. By definition, $\tilde{\mathcal{P}}$-decomposition induces a partition of the reaction set of the whole network. For each $i$, $C_i$ is partitioned by $P_i$ and $C_i \setminus P_i$. \qed

Proposition 3.6. The $\tilde{\mathcal{P}}$-decomposition is a refinement of the $\mathcal{O}$-decomposition.

Proof. This follows from the fact that $\tilde{\mathcal{O}}$-decomposition induces a partition of the reaction set of the whole network. \qed

Proposition 3.7. If a CRN is composed of irreversible reactions only, then

i. $\tilde{\mathcal{P}}$-decomposition is independent if and only if $\mathcal{F}$-decomposition is independent;

ii. $\tilde{\mathcal{P}}$-decomposition is incidence-independent if and only if $\mathcal{F}$-decomposition is incidence-independent; and,
iii. $\mathcal{P}$-decomposition is bi-independent if and only if $\mathcal{F}$-decomposition is bi-independent.

We illustrate the identified decompositions using the embedded representation of the S-system model of anti-inflammatory signaling in macrophages is given in the supplementary materials in [2].

Example 3.8. The reaction network of the embedded representation of the S-system model of anti-inflammatory signaling in macrophages is given in the supplementary materials in [2] and was referred to as CPA3-S. The biological model was based on [14]. The network is given by the following reactions.

\[ \begin{align*}
R_1 & : X_6 \to X_6 + X_1 \\
R_2 & : X_1 \to 0 \\
R_3 & : X_6 \to X_6 + X_2 \\
R_4 & : X_2 \to 0 \\
R_5 & : X_1 + X_2 + X_7 \to X_1 + X_2 + X_7 + X_3 \\
R_6 & : X_3 \to 0 \\
R_7 & : X_3 \to X_3 + X_4 \\
R_8 & : X_4 \to 0 \\
R_9 & : X_4 \to X_4 + X_5 \\
R_{10} & : X_4 + X_5 \to X_4 \\
R_{11} & : X_4 + X_5 \to X_4 + X_5 + X_6 \\
R_{12} & : X_6 \to 0 \\
R_{13} & : X_9 \to X_9 + X_7 \\
R_{14} & : X_7 + X_8 \to X_8 \\
R_{15} & : X_9 \to X_9 + X_8 \\
R_{16} & : X_7 + X_8 \to X_7 \\
R_{17} & : 0 \to X_9 \\
R_{18} & : X_6 + X_7 + X_9 \to X_6 + X_7
\end{align*} \]

The only reversible reactions are $R_9$ and $R_{10}$. We choose the orientation $\mathcal{O} = \mathcal{R} \setminus \{R_{10}\}$. The $\mathcal{O}$-decomposition is induced by the partition $\{\mathcal{R} \setminus \{R_{10}\}, \{R_{10}\}\}$.

To classify the reactions into equivalence classes, we get a basis for $\ker L_{\mathcal{O}}$ by solving the following equation.

\[ \begin{align*}
\alpha_1 X_1 + \alpha_2 (-X_1) + \alpha_3 X_2 + \alpha_4 (-X_2) + \alpha_5 X_3 + \alpha_6 (-X_3) + \alpha_7 X_4 + \alpha_8 (-X_4)
+ \alpha_9 X_5 + \alpha_{10} X_6 + \alpha_{11} (-X_6) + \alpha_{12} X_7 + \alpha_{13} (-X_7) + \alpha_{14} X_8 + \alpha_{15} (-X_8)
\end{align*} \]
\[ +\alpha_{16} X_9 + \alpha_{17} (-X_9) = 0 \]

The above equation leads to the following.
\[
\begin{align*}
\alpha_1 - \alpha_2 &= 0 \\
\alpha_3 - \alpha_4 &= 0 \\
\alpha_5 - \alpha_6 &= 0 \\
\alpha_7 - \alpha_8 &= 0 \\
\alpha_9 &= 0 \\
\alpha_{10} - \alpha_{11} &= 0 \\
\alpha_{12} - \alpha_{13} &= 0 \\
\alpha_{14} - \alpha_{15} &= 0 \\
\alpha_{16} - \alpha_{17} &= 0 
\end{align*}
\]

Hence the corresponding matrix is given by
\[
\begin{pmatrix}
R_1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
R_2 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
R_3 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
R_4 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
R_5 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
R_6 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
R_7 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
R_8 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
R_9 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
R_{11} & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
R_{12} & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
R_{13} & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
R_{14} & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
R_{15} & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
R_{16} & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
R_{17} & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
R_{18} & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1
\end{pmatrix}
\]

The equivalence classes are given by
\[
\begin{align*}
P_0 &= \{ R_9 \} \\
P_1 &= \{ R_1, R_2 \} \\
P_2 &= \{ R_3, R_4 \} \\
P_3 &= \{ R_5, R_6 \} \\
P_4 &= \{ R_7, R_8 \} \\
P_5 &= \{ R_{11}, R_{12} \} \\
P_6 &= \{ R_{13}, R_{14} \} \\
P_7 &= \{ R_{15}, R_{16} \}
\end{align*}
\]
$P_8 = \{R_{17}, R_{18}\}$

which induces the $\mathcal{P}$-decomposition.

The fundamental classes are given by

$C_0 = \{R_9, R_{10}\}$

$C_1 = \{R_1, R_2\}$

$C_2 = \{R_3, R_4\}$

$C_3 = \{R_5, R_6\}$

$C_4 = \{R_7, R_8\}$

$C_5 = \{R_{11}, R_{12}\}$

$C_6 = \{R_{13}, R_{14}\}$

$C_7 = \{R_{15}, R_{16}\}$

$C_8 = \{R_{17}, R_{18}\}$

which induces the $\mathcal{F}$-decomposition.

Now,

$P_0 = \{R_0\}$

$P_1 = \{R_1, R_2\}$

$P_2 = \{R_3, R_4\}$

$P_3 = \{R_5, R_6\}$

$P_4 = \{R_7, R_8\}$

$P_5 = \{R_{11}, R_{12}\}$

$P_6 = \{R_{13}, R_{14}\}$

$P_7 = \{R_{15}, R_{16}\}$

$P_8 = \{R_{17}, R_{18}\}$

$C_0 \setminus P_0 = \{R_{10}\}$

which induces the $\widetilde{\mathcal{P}}$-decomposition.

It is easy to check that the stoichiometric subspace of whole network is equal to the direct sum of the stoichiometric subspaces of the subnetworks under the $\mathcal{F}$-decomposition. Thus, the $\mathcal{F}$-decomposition is independent. Moreover, the rank of the of the incidence matrix is 17 which is equal to the sum of the ranks of the incidence matrices under the $\mathcal{F}$-decomposition. Thus, the $\mathcal{F}$-decomposition is incidence-independent. It follows that $\mathcal{F}$-decomposition is bi-independent.
Table 3.1. A Summary of Basic Properties of CPA3-S with respect to the $\mathcal{F}$-decomposition

| Subnetworks                                      | $n$ | $\ell$ | $s$ | $\delta$ |
|-------------------------------------------------|-----|--------|-----|----------|
| $X_6 \to X_6 + X_1, X_1 \to 0$                  | 4   | 2      | 1   | 1        |
| $X_6 \to X_6 + X_2, X_2 \to 0$                  | 4   | 2      | 1   | 1        |
| $X_1 + X_2 + X_7 \to X_1 + X_2 + X_7 + X_3, X_3 \to 0$ | 4   | 2      | 1   | 1        |
| $X_3 \to X_3 + X_4, X_4 \to 0$                  | 4   | 2      | 1   | 1        |
| $X_4 \to X_4 + X_5, X_4 + X_5 \to X_4$          | 2   | 1      | 1   | 0        |
| $X_4 + X_5 \to X_4 + X_5 + X_6, X_6 \to 0$      | 4   | 2      | 1   | 1        |
| $X_9 \to X_9 + X_7, X_7 + X_8 \to X_8$          | 4   | 2      | 1   | 1        |
| $X_9 \to X_9 + X_8, X_7 + X_8 \to X_7$          | 4   | 2      | 1   | 1        |
| $0 \to X_9, X_6 + X_7 + X_9 \to X_6 + X_7$      | 4   | 2      | 1   | 1        |

4 On Types of Fundamental Decompositions of CRNs

We begin this section with the classification of the subnetworks occurring in a $\mathcal{P}$-decomposition of a network into 3 types. Note that the subnetworks from the decomposition have deficiency either 0 or 1.

**Lemma 4.1.** [12] Let $\mathcal{N} = (\mathcal{I}, \mathcal{O}, \mathcal{R})$ be a CRN and $\mathcal{O}$ be an orientation. Let $\mathcal{N}_{\mathcal{O},i}$ for $i = 0, 1, 2, ..., w$ be defined as the subnetwork generated by all reactions in $P_i$. Then one of the following holds:

i. The reaction vectors for $\mathcal{N}_{\mathcal{O},i}$ are linearly independent, and the subnetwork $\mathcal{N}_{\mathcal{O},i}$ based on $P_i$ forms a forest (i.e., a graph with no cycle) with deficiency 0.

ii. The reaction vectors are minimally dependent, and the subnetwork $\mathcal{N}_{\mathcal{O},i}$ based on $P_i$ forms a forest with deficiency 1.

iii. The reaction vectors are minimally dependent, and the subnetwork $\mathcal{N}_{\mathcal{O},i}$ based on $P_i$ forms a big cycle (with three vertices) with deficiency 0.

We denote the subnetwork classes in i, ii, and iii of Lemma 4.1 as Type I, Type II and Type III subnetworks respectively. The classification was extended as follows: an $\mathcal{F}$-subnetwork is of type I, II or III if it contains a $\mathcal{P}$-subnetwork of type I, II or III, respectively. We denote the numbers of subnetworks (i.e., fundamental classes) for Types I, II and III with the symbols $w_I$, $w_{II}$ and $w_{III}$, respectively. We then introduce the following definition [11].
**Definition 4.2.** An $\mathcal{F}$-decomposition is said to be

i. **Type I** if it contains Type I subnetwork only.

ii. **Type II** if it contains Type II subnetwork only.

iii. **Type III** if it contains Type III subnetwork only.

The following results show that the independence, incidence-independence, and hence, bi-independence of the $\mathcal{F}$-decomposition of a CRN depend on the relationship between the deficiency of the network and the number of Type II subnetworks.

**Proposition 4.3.** [11] If a CRN has independent $\mathcal{F}$-decomposition, then $\delta \leq w_{II}$.

**Corollary 4.4.** If the deficiency $\delta > w_{II}$ of a CRN, then the $\mathcal{F}$-decomposition is not independent.

**Proposition 4.5.** If a CRN has incidence-independent $\mathcal{F}$-decomposition, then $\delta \geq w_{II}$.

**Proof.** Since the CRN has incidence-independent $\mathcal{F}$-decomposition, so $\delta \geq \delta_1 + \delta_2 + ... + \delta_w$. Thus, $\delta \geq \delta_1 + \delta_2 + ... + \delta_w$. Since each Type I or Type III subnetwork has zero deficiency, the deficiency is dependent on the number of Type II subnetworks, i.e., $\delta \geq w_{II}$. $\square$

**Corollary 4.6.** If the deficiency $\delta < w_{II}$ of a CRN, then the $\mathcal{F}$-decomposition is not incidence-independent.

**Corollary 4.7.** If a CRN has bi-independent $\mathcal{F}$-decomposition, then $\delta = w_{II}$.

**Proof.** The conclusion follows directly from Proposition 4.3 and Proposition 4.5. $\square$

**Example 4.8.** It was shown in [11] that the following CRN for $k$-site distributive phosphorylation/ dephosphorylation:

\[
S_0 + K \rightleftharpoons S_0K \rightarrow S_1 + K \rightleftharpoons S_1K \rightarrow S_2 + K \rightleftharpoons ... \rightarrow S_k + K \\
S_k + F \rightleftharpoons ... \rightarrow S_2 + F \rightleftharpoons S_2F \rightarrow S_1 + F \rightleftharpoons S_1F \rightarrow S_0 + F
\]

in [3] has bi-independent $\mathcal{F}$-decomposition with the following subnetworks:

\[
S_i + K \rightleftharpoons S_iK \rightarrow S_{i+1} + K \\
S_{i+1} + F \rightleftharpoons S_{i+1}F \rightarrow S_i + F \quad \text{for} \quad i = 0, 1, ..., k - 1.
\]

The CRN has $4k + 2$ complexes and there are 2 linkage classes. In addition, the rank of the network is $3k$. Hence, the deficiency of the CRN is $\delta = (4k + 2) - 2 - 3k = k$. Note that each of the $k$ subnetwork is of Type II (forest of deficiency 1).
We now give an example that determines whether a CRN has the capacity for multistationarity using the MATLAB program (in Appendix B) that we developed for fundamental decompositions, the DZT, and the FDT.

Example 4.9. We consider the following CRN.

\[ R_1 : M_1 \rightarrow M_2 \]
\[ R_2 : M_2 \rightarrow M_3 \]
\[ R_3 : M_3 \rightarrow M_1 \]
\[ R_4 : M_2 \rightarrow M_4 \]
\[ R_5 : M_4 \rightarrow M_5 \]
\[ R_6 : M_5 \rightarrow M_2 \]

We use the program in Appendix B with the following input.

```matlab
model.id = 'SAMPLE';
model.name = 'SAMPLE';
model.species = struct('id', {'M1', 'M2', 'M3', 'M4', 'M5'});
model.reaction(1) = struct('id', 'M1 -> M2', 'reactant', struct('species', {'M1'}, 'stoichiometry', {1}), 'product', struct('species', {'M2'}, 'stoichiometry', {1}), 'reversible', false);
model.reaction(2) = struct('id', 'M2 -> M3', 'reactant', struct('species', {'M2'}, 'stoichiometry', {1}), 'product', struct('species', {'M3'}, 'stoichiometry', {1}), 'reversible', false);
model.reaction(3) = struct('id', 'M3 -> M1', 'reactant', struct('species', {'M3'}, 'stoichiometry', {1}), 'product', struct('species', {'M1'}, 'stoichiometry', {1}), 'reversible', false);
model.reaction(4) = struct('id', 'M2 -> M4', 'reactant', struct('species', {'M2'}, 'stoichiometry', {1}), 'product', struct('species', {'M4'}, 'stoichiometry', {1}), 'reversible', false);
model.reaction(5) = struct('id', 'M4 -> M5', 'reactant', struct('species', {'M4'}, 'stoichiometry', {1}), 'product', struct('species', {'M5'}, 'stoichiometry', {1}), 'reversible', false);
model.reaction(6) = struct('id', 'M5 -> M2', 'reactant', struct('species', {'M5'}, 'stoichiometry', {1}), 'product', struct('species', {'M2'}, 'stoichiometry', {1}), 'reversible', false);
```

Then the following output is obtained.

**SUBNETWORK 1:**

\[ M_2 \rightarrow M_4 \]
\[ M_4 \rightarrow M_5 \]
\[ M_5 \rightarrow M_2 \]

**SUBNETWORK 2:**

\[ M_1 \rightarrow M_2 \]
\[ M_2 \rightarrow M_3 \]
\[ M_3 \rightarrow M_1 \]

**CONCLUSION 1:** The F-decomposition is INDEPENDENT.

**CONCLUSION 2:** The F-decomposition is INCIDENCE-INDEPENDENT.

**CONCLUSION 3:** The F-decomposition is BI-INDEPENDENT.

The subnetworks are cycles (of deficiency 0). By the DZT, each subnetwork does not have the capacity for multistationarity. By the FDT, the network cannot admit multistationarity.
The subnetwork of the Schmitz’s carbon cycle model in \cite{8,15} was generalized in \cite{10}, which is an instance of an independent Type III $F$-decomposition. An illustration of the graph is given in Figure 1.

Theorem 4.10. \cite{11} The following family of CRNs has bi-independent Type III $F$-decomposition such that the $\mathcal{N}_i$’s are precisely the fundamental classes under the decomposition: $\mathcal{N} = \{\mathcal{N}_i | \mathcal{N}_i = (\mathcal{C}_i, \mathcal{R}_i)\}$ with a (possibly broken) chain of long monomolecular directed cycles, i.e. of length $\geq 3$, and $|\mathcal{C}_i \cap \mathcal{C}_j| \leq 1$ if $j = i + 1$ for $i = 0, 1, \ldots, k - 1$.

The following result analyzes whether the mass action kinetics with underlying CRN in Theorem 4.10 has the capacity for multistationarity.

Theorem 4.11. The CRN $\mathcal{N} = \{\mathcal{N}_i | \mathcal{N}_i = (\mathcal{C}_i, \mathcal{R}_i)\}$ with a (possibly broken) chain of long monomolecular directed cycles, i.e. of length $\geq 3$, and $|\mathcal{C}_i \cap \mathcal{C}_j| \leq 1$ if $j = i + 1$ for $i = 0, 1, \ldots, k - 1$, endowed with mass action kinetics, does not have the capacity for multistationarity, i.e., the system admits at most one equilibrium.

Proof. The fundamental classes of the network under the decomposition are weakly reversible cycles of deficiency 0. By the DZT (Statement iii), each subnetwork does not have the capacity for multistationarity. By the FDT, the whole network does not have the capacity as well. \hfill $\square$

5 Fundamental Decompositions of CRNs under the CF-RM Transformation

In this section, we present a transformation method whose key property is that it maps an irreversible reaction (a reversible pair of reactions) of the original system to an irreversible reaction (a reversible pair of reactions) of the target system. In other words, it is...
reversibility and irreversibility (RI) preserving. This method was based on the generic CF-RM method (transformation of complex factorizable kinetics by reactant multiples) which converts a PL-NDK to a PL-RDK system. We add in the notation CF-RI a sub-index “+” for two reasons: to indicate the “positive” (or preserving) relation and to highlight its partial coincidence with the CF-RM$_+$ variant of CF-RM. The following presents the steps of the CF-RM transformation method in [13].

1. Determine the set of reactant complexes $\rho(\mathcal{R})$.

2. Leave each CF-reactant complex unchanged.

3. At an NF-reactant complex, select a CF-subset containing the highest number of reactions and leave this CF-subset unchanged. For each of the remaining $N_R(y) - 1$ CF-subsets, choose successively a multiple of $y$ which is not among the current set of reactants. Different procedures are possible for the selection of a new reactant as long as it is different from those in the current reactant set. After each choice, the current set is updated.

CF-RM$_+$ is a variant of CF-RM. All the steps are identical with the generic CF-RM method except that it uses additional criteria in the selection of the new reactant multiples. CF-RM$_+$ chooses the reactant multiple so that the new reactant differs from all existing complexes and all the new product complexes in the CF-subset also differ from all existing complexes [13]. Note that the CF-RM$_+$ method given in [13] updates the set of current complexes and complexes in the transform after each CF-subset of an NF-node is processed. The CF-RI$_+$ method proceeds as follows:

1. Determine the reactant set $\rho(\mathcal{R})$ and identify the subset $\rho(\mathcal{R})_{CF}$ of CF-nodes.

2. If the reaction set $\mathcal{R}_y := \rho^{-1}(y)$ of a CF-node $y$ has no reversible reaction with an NF-node, then it is left unchanged.

3. At an NF-node without reversible reactions, carry out the steps of CF-RM$_+$.

4. At an NF-node with a reversible reaction, among the CF-subsets without a reversible reaction (if there are any), select one with the highest number of reactions and leave this unchanged.

5. For the remaining CF-subsets without a reversible reaction, carry out CF-RM$_+$. 
6. For a CF-subset with a reversible reaction, carry out CF-RM\textsubscript{+}, but in addition, for each reversible reaction, also for the CF-subset of the reverse reaction (with the same “catalytic” complex). If the reactant complex of the reverse reaction is an NF-node, this additional step removes the original CF-subset from the reaction set of that NF-node. If this removal transforms the NF-node to a CF-node, then remove the node from the list of NF-nodes.

The following theorem relates the independence of the fundamental decomposition of a system and its CF-RI\textsubscript{+} transform.

**Theorem 5.1.** [11] Let \((\mathcal{N}, K)\) be a PL-NDK system and \((\mathcal{N}_\text{RI}, K_\text{RI})\) a CF-RI\textsubscript{+} transform. Then

i. for any orientation \(O\) of \(\mathcal{N}\), \(|O| = |O_{RI}|\), and

ii. the \(F\)-decomposition of \(\mathcal{N}\) is independent if and only if the \(F\)-decomposition of \(\mathcal{N}_{RI}\) is independent.

The following theorem is a restatement of Theorems 3.4 and 5.1. This implies that with or without the application of the CF-RM Transformation, the computation in the Higher Deficiency Algorithm are the same with the assumption that the \(P\)-decomposition or the \(F\)-decomposition is independent. We assume that if we have the symbol * in the notation, we are dealing with the CF-RM applied to it.

**Theorem 5.2.** Let \((\mathcal{I}, \mathcal{C}, \mathcal{R})\) be a CRN and \(O\) be an orientation. Suppose under the CF-RM transformation, the reversibility and irreversibility of the reactions are retained, with an induced orientation \(O^*\). Then \(|O| = |O^*|\) and the following are equivalent.

i. The \(P\)-decomposition is independent.

ii. The \(F\)-decomposition is independent.

iii. The \(P^*\)-decomposition is independent.

iv. The \(F^*\)-decomposition is independent.

*Proof. (i) \(\Leftrightarrow\) (iii) follows from Theorem 5.1 (i) \(\Leftrightarrow\) (ii) follows from Theorem 3.4 (iii) \(\Leftrightarrow\) (iv) follows from Theorem 3.4 Finally, (ii) \(\Leftrightarrow\) (iv) follows from Theorem 5.1. \(\Box\)*
Example 5.3. We again consider the reaction network of the embedded representation of the S-system model of anti-inflammatory signaling in macrophages in Example 3.8. Based on [2], the NF-reactant complexes are $X_9$ and $X_6$. The branching reactions at $X_9$ are $R_{13}$ and $R_{15}$. We refer to the set of differential equations in [14] regarding the kinetic order values. We choose to modify $R_{15}$: $X_9 \rightarrow X_9 + X_8$ to $R_{15}^\ast: 2X_9 \rightarrow 2X_9 + X_8$. Now, the branching reactions at $X_6$ are $R_1$, $R_3$ and $R_{12}$. In a similar manner, we choose to modify $R_{12}$: $X_6 \rightarrow 0$ to $R_{12}^\ast: 2X_6 \rightarrow X_6$. The transformation yields the following reactions.

\[
\begin{align*}
R_1 & : X_6 \rightarrow X_6 + X_1 \\
R_2 & : X_1 \rightarrow 0 \\
R_3 & : X_6 \rightarrow X_6 + X_2 \\
R_4 & : X_2 \rightarrow 0 \\
R_5 & : X_1 + X_2 + X_7 \rightarrow X_1 + X_2 + X_7 + X_3 \\
R_6 & : X_3 \rightarrow 0 \\
R_7 & : X_3 \rightarrow X_3 + X_4 \\
R_8 & : X_4 \rightarrow 0 \\
R_9 & : X_4 \rightarrow X_4 + X_5 \\
R_{10} & : X_4 + X_5 \rightarrow X_4 \\
R_{11} & : X_4 + X_5 \rightarrow X_4 + X_5 + X_6 \\
R_{12}^\ast & : 2X_6 \rightarrow X_6 \\
R_{13} & : X_9 \rightarrow X_9 + X_7 \\
R_{14} & : X_7 + X_8 \rightarrow X_8 \\
R_{15}^\ast & : 2X_9 \rightarrow 2X_9 + X_8 \\
R_{16} & : X_7 + X_8 \rightarrow X_7 \\
R_{17} & : 0 \rightarrow X_9 \\
R_{18} & : X_6 + X_7 + X_9 \rightarrow X_6 + X_7
\end{align*}
\]
Hence a basis for $\text{Ker}L_{\sigma}$ is given below.

$$
\begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 1
\end{pmatrix}
$$

This is precisely the given basis for $\text{Ker}L_{\sigma}$. Thus, the partition of $\mathcal{N}^*_{\sigma}$ into equivalence classes and $\mathcal{N}^*$ into fundamental classes yields exactly the same as that of Example 3.8. And since the reaction vectors are unchanged after the application of CF-RM, both the $\mathcal{P}^*$- and the $\mathcal{F}^*$-decompositions are also independent. This is consistent with Theorem 5.2. We can also check that both the $\mathcal{P}^*$- and the $\mathcal{F}^*$-decompositions are incidence-independent, and hence, bi-independent.

**Remark 5.4.** If reversibility and irreversibility of the reactions under the CF-RM are retained, Theorem 5.2 implies that the computation in the HDA are just the same for independent $\mathcal{P}$-decomposition or $\mathcal{F}$-decomposition. For the case where some irreversible reactions become reversible under the CF-RM, one can use Proposition 5.1. In this case, computation in the HDA might differ in the shelving assignment (you may see [10, 12] for details) since there are some irreversible reactions which became reversible under the CF-RM.

The following example will show that we do not have equivalence statements as in Theorem 5.2 for incidence-independence assuming the same conditions.

**Example 5.5.** Consider the following reaction network given in [10].

$R_1 : 0 \rightarrow A_1$

$R_2 : A_1 \rightarrow 0$
The network has a unique linkage class which is precisely a terminal strong linkage class. The rank of the network is 1 and has a deficiency of 1. Moreover, it is weakly reversible. Consider the following kinetic order values: for \( R_1: 0 \), for \( R_2: 0.5 \), for \( R_3: 1 \), and for \( R_4: 0.5 \). Thus, the system is PL-NDK. Using the CF-RM transformation, we modify \( R_3: 3A_1 \rightarrow 4A_1 \). We obtain the following network with deficiency 2 [10].

Note that the reversibility and the irreversibility of the reactions remain the same after the application of CF-RM. We can actually verify that indeed without using the CF-RM, one can directly apply the HDA and the computation yields the same results.

**Example 5.6.** Consider the following reaction network with its kinetic order matrix.

\[
\begin{pmatrix}
A & B & C \\
R_1: A \rightarrow B & 1 & 0 & 0 \\
R_2: B \rightarrow C & 0 & 1 & 0 \\
R_3: A \rightarrow 0 & 0.5 & 0 & 0 \\
R_4: B \rightarrow 0 & 0 & 0.5 & 0 \\
R_5: C \rightarrow 0 & 0 & 0 & 1 \\
\end{pmatrix}
\]

The following is a basis for \( \text{Ker}L_\sigma \).

\[
\begin{pmatrix}
1 & 1 \\
1 & 0 \\
-1 & -1 \\
0 & 1 \\
1 & 0 \\
\end{pmatrix}
\]

Consequently, the equivalence classes coincide with the fundamental classes. \( P_1 = C_1 = \{R_1, R_3\} \), \( P_2 = C_2 = \{R_2, R_5\} \), \( P_3 = C_3 = \{R_4\} \)

Below are the incidence matrices of the equivalence classes (which are equal to the corresponding fundamental classes).

\[
\begin{pmatrix}
R_1 & R_3 \\
A & -1 & 0 \\
B & 1 & -1 \\
0 & 0 & 1 \\
\end{pmatrix}
\]

\[
\begin{pmatrix}
R_2 & R_5 \\
B & -1 & 0 \\
C & 1 & -1 \\
0 & 0 & 1 \\
\end{pmatrix}
\]

\[
\begin{pmatrix}
R_4 \\
B & -1 \\
& 0 & 0 & 1 \\
\end{pmatrix}
\]

Hence, the direct sum of the incidence matrices has rank 5. On the other hand, below is
the incidence matrix of the whole network.

\[
\begin{pmatrix}
R_1 & R_2 & R_3 & R_4 & R_5 \\
A & -1 & 0 & -1 & 0 & 0 \\
B & 1 & -1 & 0 & -1 & 0 \\
C & 0 & 1 & 0 & 0 & -1 \\
0 & 0 & 0 & 1 & 1 & 1 \\
\end{pmatrix}
\]

We see that the incidence matrix has rank less than 5. Therefore, the $\mathcal{P}$-decomposition is not incidence-independent. Also, the $\mathcal{F}$-decomposition is not incidence-independent.

Now, we use the CF-RM to transform the PL-NDK to a dynamically equivalent PL-RDK. The branching nodes are $A$ and $B$. For the node $A$, the branching reactions are $R_1$ and $R_3$. We choose to change $R_3 : A \rightarrow 0$ to $R_3^* : 2A \rightarrow A$. For the node $B$, the branching reactions are $R_2$ and $R_4$. We choose to change $R_4 : B \rightarrow 0$ to $R_4^* : 2B \rightarrow B$. Hence, we have the following resulting reaction network.

\[
\begin{align*}
A & \quad B & \quad C \\
R_1 : A \rightarrow B & \quad 1 & \quad 0 & \quad 0 \\
R_2 : B \rightarrow C & \quad 0 & \quad 1 & \quad 0 \\
R_3^* : 2A \rightarrow A & \quad 0.5 & \quad 0 & \quad 0 \\
R_4^* : 2B \rightarrow B & \quad 0 & \quad 0.5 & \quad 0 \\
R_5 : C \rightarrow 0 & \quad 0 & \quad 0 & \quad 1 \\
\end{align*}
\]

Below are the incidence matrices of the equivalence classes (which are equal to the corresponding fundamental classes).

\[
\begin{pmatrix}
R_1 & R_3^* \\
A & -1 & 1 \\
B & 1 & 0 \\
2A & 0 & -1 \\
\end{pmatrix}
\quad
\begin{pmatrix}
R_2 & R_5 \\
B & -1 & 0 \\
C & 1 & -1 \\
0 & 0 & 1 \\
\end{pmatrix}
\quad
\begin{pmatrix}
R_4^* \\
2B & -1 \\
B & 1 \\
\end{pmatrix}
\]

Hence, the direct sum of the incidence matrices has rank 5. On the other hand, below is
the incidence matrix of the whole network.

\[
\begin{pmatrix}
A & \begin{pmatrix} -1 & 0 & 1 & 0 & 0 \\ B & 1 & -1 & 0 & 1 & 0 \\ C & 0 & 1 & 0 & 0 & -1 \\ 2A & 0 & 0 & -1 & 0 & 0 \\ 2B & 0 & 0 & 0 & -1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \\
\end{pmatrix}
\end{pmatrix}
\]

Thus, the incidence matrix has rank 5. Therefore, the $\mathcal{P}$-decomposition is incidence-independent. Also, the $\mathcal{F}$-decomposition is incidence-independent.

**Remark 5.7.** Even with the restriction that the reversibility and irreversibility of the reactions under the CF-RM are retained, Example 5.6 shows that if $\mathcal{P}$-decomposition is incidence-independent, it does not follow that the $\mathcal{P}^*$-decomposition is also incidence-independent. Likewise, the incidence-independence of the $\mathcal{F}$-decomposition does not imply the incidence-independence of the $\mathcal{F}^*$-decomposition.

### 6 Conclusions and Outlook

We summarize our results and provide some direction for future research.

1. We illustrated the $\mathcal{O}$-, $\mathcal{P}$-, and $\mathcal{F}$-decompositions underlying the higher deficiency algorithm for mass action kinetics and the multistationarity algorithm for power-law kinetics. We also derived properties of these decompositions.

2. We employed the use of (1) the program that we created to determine the fundamental classes and whether or not the decomposition is independent and incidence-independent, (2) the DZT, and (3) the FDT, to provide a simple solution of multistationarity of the CRN of the generalization of a subnetwork of Schmitz’s carbon cycle model by Hernandez et al. endowed with mass action kinetics.

3. We stated equivalent statements regarding the $\mathcal{P}$-, the $\mathcal{P}^*$-, the $\mathcal{F}$-, and the $\mathcal{F}^*$-decompositions with the assumption that the reversibility and irreversibility of the reactions under the CF-RM are retained. In this case, $|\mathcal{O}| = |\mathcal{O}^*|$. 
4. Even with the restriction that the reversibility and irreversibility of the reactions under the CF-RM are preserved, Example 5.6 shows that if $\mathcal{F}$-decomposition is incidence-independent, it does not follow that the $\mathcal{F}^*$-decomposition is also incidence-independent. We provided a counterexample for this case.

5. One may look into necessary conditions to establish equivalence statements for fundamental incidence-independent decompositions.

References

[1] C. P. Arceo, E. Jose, A. Marín-Sanguino, E. Mendoza, Chemical reaction network approaches to Biochemical Systems Theory, *Math. Biosci.* **269** (2015) 135–152.

[2] C. P. Arceo, E. Jose, A. Lao, E. Mendoza, Reaction networks and kinetics of biochemical systems, *Math. Biosci.* **283** (2017) 13–29.

[3] C. Conradi, A. Shiu, Dynamics of post-translational modification systems: recent results and future directions, *Biophysical Journal* **114**(3) (2018) 507–515.

[4] H. Fariñas, E. Mendoza, A. Lao, Decompositions of chemical reaction networks and embedded networks of S-systems, in preparation.

[5] M. Feinberg, Chemical reaction network structure and the stability of complex isothermal reactors I: The deficiency zero and deficiency one theorems, *Chem. Eng. Sci.* **42** (1987) 2229–2268.

[6] M. Feinberg, Lectures on chemical reaction networks. Notes of lectures given at the Mathematics Research Center of the University of Wisconsin, 1979. Available at https://crnt.osu.edu/LecturesOnReactionNetworks.

[7] M. Feinberg, The existence and uniqueness of steady states for a class of chemical reaction networks, *Arch. Ration. Mech. Anal.* **132** (1995) 311–370.

[8] N. Fortun, A. Lao, L. Razon, E. Mendoza, A deficiency zero theorem for a class of power-law kinetic systems with non-reactant-determined interactions, *MATCH Commun. Math. Comput. Chem.* **81**(3) (2019) 621–638.

[9] E. Gross, H. Harrington, N. Meshkat, A. Shiu, Joining and decomposing reaction networks, (2018, submitted).

[10] B. Hernandez, E. Mendoza, A. de los Reyes V, A computational approach to multistationarity of power-law kinetic systems, *J Math Chem* **58**(1) (2020) 56–87. DOI: 10.1007/s10910-019-01072-7
[11] B. Hernandez, E. Mendoza, A. de los Reyes V, Fundamental decompositions and multistationarity of power-law kinetic systems, *MATCH Commun. Math. Comput. Chem.* **83**(2) (2020) 403-434.

[12] H. Ji, Uniqueness of equilibria for complex chemical reaction networks, Ph.D. Dissertation, Ohio State University, 2011.

[13] A. Nazareno, R. Eclarin, E. Mendoza, A. Lao, Linear conjugacy of chemical kinetic systems, *Math. Biosci. Eng.* **16**(6) (2019) 8322-8355.

[14] J. Ray, D. Kirschner, Requirement for multiple activation signals by anti-inflammatory feedback in macrophages, *J. Theor. Biol.* 241 (2006) 276-294.

[15] R. Schmitz, The Earth’s carbon cycle: Chemical engineering course material, *Chemical Engineering Education* **36**(4) (2002) 296–309.

[16] N. Soranzo, C. Altafini, ERNEST: a toolbox for chemical reaction network theory, *Bioinformatics* 25 (2009) 2853-2854.

A  Nomenclature

A.1  List of abbreviations

| Abbreviation | Meaning |
|--------------|---------|
| CF | complex factorizable |
| CKS | chemical kinetic system |
| CRN | chemical reaction network |
| CRNT | Chemical Reaction Network Theory |
| GMA | generalized mass action |
| HDA | higher deficiency algorithm |
| MAK | mass action kinetics |
| MSA | multistationarity algorithm |
| PLK | power-law kinetics |
| PL-NDK | power-law non-reactant-determined kinetics |
| PL-RDK | power-law reactant-determined kinetics |
| SFRF | species formation rate function |
### A.2 List of important symbols

| Meaning                                      | Symbol |
|----------------------------------------------|--------|
| deficiency                                   | δ      |
| dimension of the stoichiometric subspace     | s      |
| incidence map                                | I_{a}  |
| molecularity matrix                          | Y      |
| number of complexes                          | n      |
| number of linkage classes                    | l      |
| number of strong linkage classes             | s_{l}  |
| orientation                                  | O      |
| stoichiometric matrix                        | N      |
| stoichiometric subspace                      | S      |
| subnetwork of \( \mathcal{N} \) with respect to \( \mathcal{O} \) | \( \mathcal{N}_{\mathcal{O}} \) |

### B A MATLAB program

We now provide a MATLAB program that computes for the subnetworks under the fundamental decomposition of a reaction network. It also determines whether the decomposition is independent, incidence-independent and bi-independent. We use the preliminary steps of the program of Soranzo and Altafini [16] to come up with our own program. We should install the free software ERNEST in your MATLAB environment. The script was named incinde.m.

```matlab
function [ ret ] = HDA( model )
    % species = { model.species.id }; % number of species
    % reactions = { model.reaction.id }; % number of reactions
    reactant_complexes = []; % matrix of reactant complexes (species x irrev. reactions)
    product_complexes = []; % matrix of product complexes (species x irrev. reactions)
    rec = numel( reactions ); % number of reactions (counting reversible reactions as one)
    SnoRev = []; % stoichiometric matrix (without reverse of reversible reaction)
    P_i = []; % equivalence classes
    P_i_neu = []; % equivalence classes
    StoichMatrixForm = [];
    arr = [];
    for i = 1: rec
        if isfield( model.reaction(i), 'modifier' ) && ~isempty( model.reaction(i).modifier )
            warning( 'Reaction ' num2str(i) ' contains modifiers, which will be ignored. Specify all species in a reaction as reactants or products.' )
        end
        reactant_complexes(:, end+1) = zeros( n, 1 );
        for j = 1: rec
            reactant_complexes(find(strcmp(model.reaction(i).reactant(j).species, species)), end) = model.reaction(i).reactant(j).stoichiometry;
        end
        product_complexes(:, end+1) = zeros( n, 1 );
        for j = 1: rec
            product_complexes(find(strcmp(model.reaction(i).product(j).species, species)), end) = model.reaction(i).product(j).stoichiometry;
        end
        SnoRev(:, end + 1) = product_complexes(:, end) - reactant_complexes(:, end);
    end
    clear label
    for i = 1: rec
        if isfield( model.reaction(i), 'modifier' ) && ~isempty( model.reaction(i).modifier )
            warning( 'Reaction ' num2str(i) ' contains modifiers, which will be ignored. Specify all species in a reaction as reactants or products.' )
        end
        reactant_complexes(:, end+1) = zeros( n, 1 );
    end
```
for j = 1:numel(model.reaction(i).reactant)
    reactant_complexes(find(strcmp(model.reaction(i).reactant(j).species, species), 1), end) = model.reaction(i).reactant(j).stoichiometry;
end
product_complexes(:, end+1) = zeros(n, 1);
for j = 1:numel(model.reaction(i).product)
    product_complexes(find(strcmp(model.reaction(i).product(j).species, species), 1), end) = model.reaction(i).product(j).stoichiometry;
end
S(:, end + 1) = product_complexes(:, end) - reactant_complexes(:, end);
if model.reaction(i).reversible
    reactant_complexes(:, end+1) = product_complexes(:, end);
    product_complexes(:, end+1) = reactant_complexes(:, end-1);
    S(:, end + 1) = -S(:, end);
end
if numel(model.reaction(i).reactant) > 0 && numel(model.reaction(i).product) > 0
    label = [num2str(model.reaction(i).reactant(1).stoichiometry) ' ' model.reaction(i).reactant(1).species];
    for j = 2:numel(model.reaction(i).reactant)
        label = [label ' + ' num2str(model.reaction(i).reactant(j).stoichiometry) ' ' model.reaction(i).reactant(j).species];
    end
    sr_edges(end+1,:) = {model.reaction(i).reactant(1).species, reactions{i}, label, model.reaction(i).reactant(1).stoichiometry};
end
    label = [num2str(model.reaction(i).product(1).stoichiometry) ' ' model.reaction(i).product(1).species];
    for j = 2:numel(model.reaction(i).product)
        label = [label ' + ' num2str(model.reaction(i).product(j).stoichiometry) ' ' model.reaction(i).product(j).species];
    end
    for j = 1:numel(model.reaction(i).product)
        sr_edges(end+1,:) = {model.reaction(i).product(j).species, reactions{i}, label, model.reaction(i).product(j).stoichiometry};
    end
end
end
clear label
[Y, ind, ind2] = unique([reactant_complexes product_complexes]', 'rows'); % ind2(i) is the index in Y of the reactant complex in reaction i, ind(i + r) is the index in Y of the product complex in reaction i
Y = Y'; % complexes matrix (species x complexes)
m = size(Y, 2); % number of complexes
reacts_to = false(m, m); % matrix (complexes x complexes) for the reacts_to relation:
    react_to(i, j) = true iff i->j
r = size(reactant_complexes, 2); % number of irrev. reactions
reacts_in = zeros(m, r); % matrix (complexes x irrev. reactions) for the reacts_in relation: (reacts_in(i, r) = -1 && reacts_in(j, r) = 1) iff i->j
for i = 1:r:
    reacts_to(ind2(i), ind2(i + r)) = true;
    reacts_in(ind2(i), i) = -1;
    reacts_in(ind2(i+r), i) = 1; % incidence
end
is_reversible = isequal(reacts_to, react_to.'); % test for reversibility
complexes_uigraph_cc = connected_components(umultigraph(reacts_to | react_to')); % linkage classes
l = max(complexes_uigraph_cc); % number of linkage classes
if is_reversible
    complexes_graph_scc = complexes_uigraph_cc;
else
    complexes_graph_scc = strongly_connected_components(multigraph(reacts_to)); % strong-linkage classes
end
n_slc = max(complexes_graph_scc); % number of strong-linkage classes
is_weakly_reversible = n_slc == l; % the reaction network is weakly reversible if and only if each linkage class is a strong-linkage class
s = rank(S); % reaction network rank
d = m - 1 - s; % reaction network deficiency
Ysp = null(SnRev.'r');
[NspRow, NspCol] = size(Ysp);
NspEqual = Nsp;
for i = 1:NspRow
    if NspEqual(i,:) == zeros(NspCol,1)
        NspEqual(i,:) = NspEqual(i,:) ./ gcd(sym(NspEqual(i,:)));
    end
end
for i = 1:NspRow
    for j = 1:NspRow
        if NspEqual(i,:) == -NspEqual(j,:)
            NspEqual(j,:) = NspEqual(i,:);
        end
    end
end
NspEqual;
NspUnique = unique(Nsp, 'rows', 'stable');
TranposeNspUnique = NspUnique.';
size(NspUnique);
NspOrtho = null(TranposeNspUnique,'r');
NspOrthoTranspose = NspOrtho.';
size(NspOrthoTranspose);
SimplifiedNspOrthoTranspose = rref(NspOrthoTranspose);
TranposeSimplifiedNspOrthoTranspose = SimplifiedNspOrthoTranspose.';
size(TranposeSimplifiedNspOrthoTranspose);
TranposeNspUnique = TranposeSimplifiedNspOrthoTranspose(:,1);
size(TranposeNspUnique);
TranposeSimplifiedOrthoTranspose = TranposeSimplifiedNspOrthoTranspose(:,1);
UniqueNspMatrix = unique(NspEqual, 'rows'); % take the unique NSP complexes
[UNSPSizeRow, UNSPSizeCol] = size(UniqueNspMatrix);
[NSPSizeRow, NSPSizeCol] = size(Nsp);
% The following command is to identify the location of reactions (equivalence classes).
[LiaRR, LocbRR] = ismember(Nsp, UniqueNspMatrix, 'rows');
IdentifyLocbRR = LocbRR;
IdentifyUniLocbRR = unique(IdentifyLocbRR('ismem(IdentifyLocbRR)));
histIdentifyLocbRR = histc(IdentifyLocbRR, IdentifyUniLocbRR);
UniqueRR = IdentifyUniLocbRR(histIdentifyLocbRR >1);
[RRSizeRow, RRSizeCol] = size(UniqueRR);
% The following command is to identify the location of reactions (equivalence classes).
[Lia, Locb] = ismember(NspEqual, UniqueNspMatrix, 'rows');
IdentifyLocb = Locb;
IdentifyUniLocb = unique(IdentifyLocb('ismem(IdentifyLocb)));
histIdentifyLocb = histc(IdentifyLocb, IdentifyUniLocb);
UniqueReactionRow = IdentifyUniLocb(histIdentifyLocb >1);
[URSizeRow, URSizeCol] = size(UniqueReactionRow);
UniqueReactionRowMatrixTik = zeros(UNSPSizeRow, numel(reactions));
for k = 1:URSizeRow
    recordALL = find(ismember(IdentifyLocb, UniqueReactionRow(k)));
    [Res, LocRes] = ismember(Nsp,(zeros(NspSizeCol,1),.'), 'rows');
    not(Res);
    if not(Res) & not(ismpty(Nsp))
        record = find(ismember(IdentifyLocb, UniqueReactionRow(k)));
        [URSizeRecordRow, URSizeRecordCol] = size(record);
        for i = 1:URSizeRecordRow
            % disp(model.reaction(record(i)).id);
            % disp(model.reaction(record(i)).reversible);
        end
        else
            Res;
            [Res, LocRes] = ismember(Nsp,(zeros(NspSizeCol,1),.'), 'rows');
            if isempty(Nsp) | Res
                fprintf('The zeroth fundamental class F0 has the following reaction(s).');
                record0 = find(ismember(IdentifyLocb, UniqueReactionRow(k)));
                fprintf('
');
                [URSizeRecordRow0, URSizeRecordCol0] = size(record0);
                for i = 1:URSizeRecordRow0
                    % disp(model.reaction(record0(i)).id);
                    % disp(model.reaction(record0(i)).reversible);
                end
            else
                fprintf('The fundamental class F%d has the following reaction(s).',
                    UniqueReactionRow(k)-1);
                record = find(ismember(IdentifyLocb, UniqueReactionRow(k)));
                fprintf('
');
[URsizeRecordRow,URsizeRecordCol] = size(record);
for i = 1:URsizeRecordRow
    %disp(model.reaction(record(i)).id);
    %disp(model.reaction(record(i)).reversible);
end
end
end
SN = Locb';
Locb = [];
subnetworks = {SN};
subnetworks2 = [SN];
subnetworks3 = unique(subnetworks2);
reactant_complexes = [];
product_complexes = [];
S = [];
sum = 0;
sum2 = 0;
StoichMatrixForm = [];
sr_edges = cell(0, 4);
%NOTE: If you want the reverse reaction of a reversible reaction to be included in a different subnetwork,
%use 2 model reactions and treat the two reactions as irreversible.
[Lia, Locb] = ismember(subnetworks2, subnetworks3);
IdentifyLocb = Locb;
IdentifyUniqLocb = unique(IdentifyLocb(~ isnan(IdentifyLocb)));
histIdentifyLocb = histc(IdentifyLocb, IdentifyUniqLocb);
UniqueReactionRow = IdentifyUniqLocb(histIdentifyLocb >= 1);
[URsizeRow, URsizeCol] = size((UniqueReactionRow)');
for k = 1:URsizeRow
    record = find(ismember(IdentifyLocb, UniqueReactionRow(k)));
    [URsizeRecordRow, URsizeRecordCol] = size(record);
    fprintf(' SUBNETWORK %d:', UniqueReactionRow(k))
    fprintf('
')
    for i = 1:URsizeRecordCol
        disp(model.reaction(record(i)).id);
        if isfield(model.reaction(record(i)), 'modifier') && ~isempty(model.reaction(record(i)).modifier)
            warning([' Reaction ', num2str(record(i)) ', contains modifiers, which will be ignored. Specify all species in a reaction as reactants or products. '])
        end
    end
    reactant_complexes(:, end + 1) = zeros(n, 1);
    for j = 1: numel(model.reaction(record(i)).reactant)
        reactant_complexes(find(strcmp(model.reaction(record(i)).reactant(j).species, species)), 1) = model.reaction(record(i)).reactant(j).stoichiometry;
    end
    product_complexes(:, end + 1) = zeros(n, 1);
    for j = 1: numel(model.reaction(record(i)).product)
        product_complexes(find(strcmp(model.reaction(record(i)).product(j).species, species)), 1) = model.reaction(record(i)).product(j).stoichiometry;
    end
    S(:, end + 1) = product_complexes(:, end) - reactant_complexes(:, end);
    if model.reaction(record(i)).reversible
        reactant_complexes(:, end + 1) = product_complexes(:, end);
        product_complexes(:, end + 1) = reactant_complexes(:, end - 1);
        S(:, end + 1) = -S(:, end);
    end
    if numel(model.reaction(record(i)).reactant) > 0 && numel(model.reaction(record(i)).product) > 0
        label = [num2str(model.reaction(record(i)).reactant(1).stoichiometry) ' ', model.reaction(record(i)).reactant(1).species];
        for j = 2: numel(model.reaction(record(i)).reactant)
            label = [label ' + ' num2str(model.reaction(record(i)).reactant(j).stoichiometry) ' ', model.reaction(record(i)).reactant(j).species];
        end
        for j = 1: numel(model.reaction(record(i)).reactant)
            sr_edges(end + 1, :) = (model.reaction(record(i)).reactant(j).species, reactions(record(i)), label, model.reaction(record(i)).reactant(j).stoichiometry);
end

label = [num2str(model.reaction(record(i)).product(1).stoichiometry) \ ' ' model.
reaction(record(i)).product(1).species];
for j = 2:numel(model.reaction(record(i)).product)
  label = [label \ '+ ' num2str(model.reaction(record(i)).product(j).
stoichiometry) \ ' ' model.reaction(record(i)).product(j).species];
end
for j = 1:numel(model.reaction(record(i)).product)
sr_edges(end+1,:) = {model.reaction(record(i)).product(j).species, reactions
(record(i)), label, model.reaction(record(i)).product(j).stoichiometry};
end
clear label

[Y, ind, ind2] = unique([reactant_complexes product_complexes]', 'rows'); % ind2(i) is
the index in Y of the reactant complex in reaction i, ind(i+r) is the index in Y of
the product complex in reaction i
Y = Y'; % complexes matrix (species x complexes)

m = size(Y, 2); % number of complexes
reacts_to = false(m, m); % matrix (complexes x complexes) for the reacts_to relation:
  reacts_to(i, j) = true iff i->j
r = size(reactant_complexes, 2); % number of irrev. reactions
reacts_in = zeros(m, r); % matrix (complexes x irrev. reactions) for the reacts_in
relation: (reacts_in(i, r) = -1 & reacts_in(j, r) = 1) iff i->j
for i = 1:r
  reacts_to(ind2(i), ind2(i+r)) = true;
  reacts_in(ind2(i), i) = -1;
  reacts_in(ind2(i+r), i) = 1; % incidence
end

is_reversible = isequal(reacts_to, reacts_to'); % test for reversibility
complexes_ugraph_cc = connected_components((umultigraph(reacts_to | reacts_to'))); %
linkage classes
l = max(complexes_ugraph_cc); % number of linkage classes
if is_reversible
  complexes_graph_scc = complexes_ugraph_cc;
else
  complexes_graph_scc = strongly_connected_components((multigraph(reacts_to)));
end

n_slc = max(complexes_graph_scc); % number of strong-linkage classes
is_weakly_reversible = n_slc == l; % the reaction network is weakly reversible if and
only if each linkage class is a strong-linkage class
s = rank(S); % reaction network rank
rr = numel(reactions); % number of reactions (counting reversible reactions as one)

%fprint('The stoichiometric subspace of SUBNETWORK %d is:', UniqueReactionRow(k))
S;
fprint('The rank of SUBNETWORK %d is:', UniqueReactionRow(k))
s = rank(S);
m;
l;
%fprint('The value of (n-1) for the SUBNETWORK %d is:', UniqueReactionRow(k))
diff=m-1;
fprint('The value of (n-1) for the SUBNETWORK %d is:', UniqueReactionRow(k))
arr=[arr; UniqueReactionRow(k) s diff];
S = {}; reactant_complexes = {}; product_complexes = {}; sum = sum + s;
sum2 = sum2 + diff;
end

%fprint('Summary of the the values of s (2ND COL) and n-l (3RD COL) of SUBNETWORK i:');
summary=[];
summary=arr;
summary
%fprint('The SUM of the RANKS of the SUBNETWORKS is:')
s=0;
%fprint('The SUM of the values of (n-l) of the SUBNETWORKS is:')
sum=
sum2
S=[];
for i = 1:numel(reactions)
    if isfield(model.reaction(i), 'modifier') && ~isempty(model.reaction(i).modifier)
        warning(['Reaction ', num2str(i), ' contains modifiers, which will be ignored. Specify all species in a reaction as reactants or products.'])
    end
    reactant_complexes(:, end+1) = zeros(n, 1);
    for j = 1:numel(model.reaction(i).reactant)
        reactant_complexes(find(strcmp(model.reaction(i).reactant(j).species, species), 1), end) = model.reaction(i).reactant(j).stoichiometry;
    end
    product_complexes(:, end+1) = zeros(n, 1);
    for j = 1:numel(model.reaction(i).product)
        product_complexes(find(strcmp(model.reaction(i).product(j).species, species), 1), end) = model.reaction(i).product(j).stoichiometry;
    end
    S(:, end + 1) = product_complexes(:, end) - reactant_complexes(:, end);
    if model.reaction(i).reversible
        reactant_complexes(:, end+1) = product_complexes(:, end);
        product_complexes(:, end+1) = reactant_complexes(:, end-1);
        S(:, end + 1) = -S(:, end);
    end
    if numel(model.reaction(i).reactant) > 0 && numel(model.reaction(i).product) > 0
        label = [num2str(model.reaction(i).reactant(1).stoichiometry) ' ' model.reaction(i).reactant(1).species];
        for j = 2:numel(model.reaction(i).reactant)
            label = [label ' + ' num2str(model.reaction(i).reactant(j).stoichiometry) ' ' model.reaction(i).reactant(j).species];
        end
        for j = 1:numel(model.reaction(i).reactant)
            sr_edges(end+1,:) = {model.reaction(i).reactant(j).species, reactions{i}, label, model.reaction(i).reactant(j).stoichiometry};
        end
        label = [num2str(model.reaction(i).product(1).stoichiometry) ' ' model.reaction(i).product(1).species];
        for j = 2:numel(model.reaction(i).product)
            label = [label ' + ' num2str(model.reaction(i).product(j).stoichiometry) ' ' model.reaction(i).product(j).species];
        end
        for j = 1:numel(model.reaction(i).product)
            sr_edges(end+1,:) = {model.reaction(i).product(j).species, reactions{i}, label, model.reaction(i).product(j).stoichiometry};
        end
    end
end

fprintf(' The stoichiometric subspace of the WHOLE NETWORK is: ')
S;
s=rank(S);
fprintf(' The rank of the WHOLE NETWORK is: ')
s
fprintf('\n')
clear label
[Y, ind, ind2] = unique([reactant_complexes product_complexes], 'rows'); % ind2(i) is the index in Y of the product complex in reaction i
Y = Y'; % complexes matrix (species x complexes)
m = size(Y, 2); % number of complexes
reacts_to = false(m, m); % matrix (complexes x complexes) for the reacts_to relation:
    reacts_to(i, j) = true iff i->j
r = size(reactant_complexes, 2); % number of irrev. reactions
reacts_in = zeros(m, r); % matrix (complexes x irrev. reactions) for the reacts_in relation:
    (reacts_in(i, r) = -1 && reacts_in(j, r) = 1) iff i->j
for i = 1:r
    reacts_to(ind2(i), ind2(i + r)) = true;
    reacts_in(ind2(i+1), i) = -1;
    reacts_in(ind2(1), i) = 1; % incidence
end
is_reversible = isequal(reacts_to, reacts_to'); % test for reversibility
complexes_ugraph_cc = connected_components(multigraph(reacts_to | reacts_to')); % linkage classes
l = max(complexes_ugraph_cc); % number of linkage classes
if is_reversible
    complexes_graph_scc = complexes_ugraph_cc;
else
    complexes_graph_scc = strongly_connected_components(multigraph(reacts_to)); \% strong-linkage classes
end
n_slc = max(complexes_graph_scc); \% number of strong-linkage classes
is_weakly_reversible = n_slc == 1; \% the reaction network is weakly reversible if and only if each linkage class is a strong-linkage class
s = rank(S); \% reaction network rank
fprintf('The value of (n-1) for the WHOLE NETWORK is: ')
diff=m-l
fprintf('NOTE: The subnetworks given above correspond to the fundamental class under the F-decomposition. ')
fprintf('
')
if sum == s
    fprintf('CONCLUSION 1: The F-decomposition is INDEPENDENT. ')
else
    fprintf('CONCLUSION 1: The F-decomposition is NOT INDEPENDENT. ')
end
fprintf('
')
if sum2 == diff
    fprintf('CONCLUSION 2: The F-decomposition is INCIDENCE-INDEPENDENT. ')
else
    fprintf('CONCLUSION 2: The F-decomposition is NOT INCIDENCE-INDEPENDENT. ')
end
fprintf('
')
if sum == s & sum2 == diff
    fprintf('CONCLUSION 3: The F-decomposition is BI-INDEPENDENT. ')
else
    fprintf('CONCLUSION 3: The F-decomposition is NOT BI-INDEPENDENT. ')
end