# Wiener number of vertex-weighted graphs and a chemical application 

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Received 15 February 1996; received in revised form 26 March 1997; accepted 31 March 1997


#### Abstract

The Wiener number $W(G)$ of a graph $G$ is the sum of distances between all pairs of vertices of $G$. If ( $G, w$ ) is a vertex-weighted graph, then the Wiener number $W(G, w)$ of $(G, w)$ is the sum, over all pairs of vertices, of products of weights of the vertices and their distance. For $G$ being a partial binary Hamming graph, a formula is given for computing $W(G, w)$ in terms of a binary Hamming labeling of $G$. This result is applied to prove that $W(P H)=W(\widetilde{H S})+36 W(I D)$, where $P H$ is a phenylene, $\widehat{H S}$ a pertinently vertex-weighted hexagonal squeeze of $P H$, and $I D$ the inner dual of the hexagonal squeeze.


## 1. Introduction

In this paper we are concerned with a graph invariant $W=W(G)$, defined for an arbitrary connected graph $G$ as

$$
W(G)=\frac{1}{2} \sum_{u, v \in V(G)} d_{G}(u, v),
$$

where $d_{G}(u, v)$ denotes the length of a shortest path in $G$ between vertices $u$ and $v$. We call $W$ the Wiener number.

The investigation of the quantity $W$ seems to be first undertaken by Harold Wiener [33] almost exactly 50 years ago. Since then there is a continuous record of research activity in this field with no sign of attenuation. Wiener's original article [33] appeared in a Chemistry journal and was long overlooked by mathematicians. In the meantimc the sum of all distances of graphs was studied in quite a few mathematical works.

[^0]A variety of names for $W$ was proposed: "gross status" [19], "total status" [4], "graph distance" [9], "transmission" [29] and simply "sum of all distances" [11, 37]. In some works the closely related "mean distance" $[8,36]$ or "average distance" $[1,6]$ were considered. The name "Wiener number" or "Wiener index" is nowadays in standard use in chemistry and is sometimes encountered also in the mathematical literature [25, 26].

In chemistry the Wiener number is one of the most thoroughly studied, best understood and most frequently used graph-theory-based molecular-shape descriptors; for reviews on chemical aspects of $W$ and further references see [18, 27]. $W$ found numerous applications in the modelling of physico-chemical, pharmacological and biological properties of organic molecules. To give an idea about the versatility of the applications of the Wiener number, we mention its use in the study of ultrasonic sound velocities in alkanes and alcohols [30], rates of electroreduction of chlorobenzenes [3], cytostatic and antihistaminic activities of certain drugs [24], protonation constants of derivatives of salicylhydroxamic acid and their fungicidal activities [22], and isomerism in fullerenes [28]. The success of $W$ was long thought to lie in its (putative) capability of measuring molecular volumes, surfaces and/or surface-to-volume ratios. It was shown quite recently [16] that, indeed, $W$ is proportional to molecular surface areas. On the other hand, contrary to earlier expectations, no relation between $W$ and molecular volumes could be established [16].

A graph $G$ is called a partial binary Hamming graph, if each vertex $v \in V(G)$ can be labelled by a word of fixed length, $\ell(v)$, defined on a two-letter alphabet, say on the alphabet $\{0,1\}$, such that for all $u, v \in V(G)$ the number of positions in which $\ell(u)$ and $\ell(v)$ differ equals $d_{G}(u, v)$, cf. [21]. The labeling $\ell$ is called a binary Hamming labeling. Partial binary Hamming graphs can also be described as isometric subgraphs of hypercubes. We refer to $[2,5,7,20,21,34,35]$ for many results and aspects of partial (binary) Hamming graphs.

In the next section we introduce the Wiener number, $W(G, w)$, of a vertex-weighted graph ( $G, w$ ), and give a formula for computing $W(G, w)$ in the case when $G$ is a partial binary Hamming graph. In Section 3 two classes of graphs of interest in chemistry are discussed: hexagonal systems and phenylenes. In particular, a distance based definition of an elementary cut of a hexagonal system is presented. Finally, in the last section, a formula is obtained for the Wiener number of a phenylene in terms of its hexagonal squeeze.

## 2. The Wiener number of vertex-weighted graphs

A vertex-weighted graph $(G, w)$ is a graph $G$ together with a function $w: V(G) \rightarrow \mathbb{N}^{+}$. (Evidently, we could have chosen for vertex-weights (positive) real numbers. For the present considerations, however, weighting of the vertices with positive integers will suffice.) From now on vertex-weighted graphs in which all weights are positive integers will be called simply weighted graphs.

The Wiener number $W(G, w)$ of a weighted graph $(G, w)$ is defined as

$$
W(G, w)=\frac{1}{2} \sum_{u, v \in V(G)} w(u) w(v) d_{G}(u, v) .
$$

Note that if $w(u)=1$ holds for all vertices $u \in V(G)$, then $W(G, w)=W(G)$. More generally, if $w$ is a constant function, say $w \equiv m$, then $W(G, w)=m^{2} W(G)$. Observe also that if the distance matrix of a graph $G$ is given, then it is no more difficult to compute $W(G, w)$ than $W(G)$.

Wiener indices of weighted graphs, as defined above, seem not to be previously studied (in either mathematical or chemical literature). Exceptionally, one of the present authors did some work [14] on $W\left(G, w^{\prime}\right)$, where the weight $w(u)$ was set to be equal to the degree of the vertex $u$.

Let ( $G, w$ ) be a weighted partial binary Hamming graph, i.e., a weighted graph ( $G, w$ ) where $G$ is a partial binary Hamming graph. Let $V(G)=\left\{v_{1}, v_{2}, \ldots, v_{n}\right\}$ and set

$$
\tilde{V}(G)=\left\{v_{1}^{1}, \ldots, v_{1}^{w(1)}, v_{2}^{1}, \ldots, v_{2}^{w(2)}, \ldots, v_{n}^{1}, \ldots, v_{n}^{w(n)}\right\},
$$

where the abbreviation $w(i)$ is used instead of $w\left(v_{i}\right)$. Let $g$ be a binary Hamming labeling of $G$. We define a labeling $\tilde{g}$ of the elements of $\tilde{V}(G)$ by $\widetilde{g}\left(v_{i}^{j}\right)=g\left(v_{i}\right)$ for all $i$ and $j$. We call $\tilde{g}$ an extended binary Hamming labeling. For $u \in \tilde{V}(G)$ let $\widetilde{g}_{i}(u)$ denote the $i$ th coordinate of the label $g(u)$.

With the above notation we have:
Proposition 1. Let $G$ be a partial binary Hamming graph with a binary Hamming labeling $g$ of length $k$. Let $\widetilde{g}$ be the extended binary Hamming labeling of the weighted partial binary Hamming graph ( $G, w$ ). Then

$$
W(G, w)=\sum_{i=1}^{k} m_{i}\left(m-m_{i}\right)
$$

where $m=\sum_{u \in V(G)} w(u)=|\dot{V}(G)|$, and $m_{i}$ is the number of vertices $u \in \dot{V}(G)$ with $\widetilde{g}_{i}(u)=1$, for $i=1,2, \ldots, k$.

Proof. Let $V=V(G)$. For $u, v \in V$, let $\delta_{i}(u, v)$ be 0 if $g_{i}(u)=g_{i}(v)$, and 1 otherwise. Since $g$ is a binary Hamming labeling, we have:

$$
\begin{aligned}
W(G, w) & =\frac{1}{2} \sum_{u \in V} \sum_{v \in V} w(u) w(v) d_{G}(u, v) \\
& =\frac{1}{2} \sum_{u \in V} \sum_{v \in V} w(u) w(v) \sum_{i=1}^{k} \delta_{i}(u, v)
\end{aligned}
$$

$$
\begin{aligned}
& =\sum_{i=1}^{k}\left(\frac{1}{2} \sum_{u \in V} \sum_{v \in V} w(u) w(v) \delta_{i}(u, v)\right) \\
& =\sum_{i=1}^{k} m_{i}\left(m-m_{i}\right) .
\end{aligned}
$$

The following special case of Proposition 1 is a previously known result [23, Proposition 3.1]:

Corollary 2. Let $G$ be a partial binary Hamming graph on $n$ vertices and let $g$ be a binary Hamming labeling of $G$ of length $k$. For $1=1,2, \ldots, k$ let $n_{i}$ be the number of vertices $u$ of $G$ with $g_{i}(u)=1$. Then

$$
W(G)=\sum_{i=1}^{k} n_{i}\left(n-n_{i}\right)
$$

Proposition 1 provides a simple method for the calculation of the Wiener numbers of (weighted) partial binary Hamming graphs, which is particularly suitable for the chemically very important class of hexagonal systems, described in the subsequent section.

## 3. Hexagonal systems and phenylenes

Hexagonal systems (or benzenoid systems or benzenoid graphs, see below) are finite connected plane graphs with no cut vertices, in which every interior region is bounded by a regular hexagon of side length 1, cf. [31, 39]. The inner dual of a hexagonal system $G$ is a graph with vertices corresponding to the hexagons of $G$, and two vertices are adjacent if the corresponding hexagons share an edge.

Hexagonal systems provide the graph representation of the so-called benzenoid hydrocarbons, a class of substances of outstanding importance in chemistry. Therefore they are often referred to as benzenoid graphs or benzenoid systems. Details of the extensive research on these molecular graphs can be found in the book [15] and the review [13]. Recall that the number of known benzenoid hydrocarbons is well above 500 ; some of them play a major role in chemical industry, some are notorious pollutants; a few of them are highly carcinogenic.

Let $G$ be a hexagonal system in which no three hexagons meet in a vertex, i.e., let $G$ be a tree-like hexagonal system. (In chemistry these are referred to as catacondensed benzenoid systems [15]). Then separate each pair of hexagons that share an edge $u v$ as follows. If $u^{\prime}, v^{\prime}$ and $u^{\prime \prime}, v^{\prime \prime}$ are the corresponding vertices in the separated cycles, join $u^{\prime}$ with $u^{\prime \prime}$ and $v^{\prime}$ with $v^{\prime \prime}$, and repeat this for all pairs of adjacent hexagons of $G$. The graph obtained in this way is called a phenylene and the graph $G$ is called


Fig. 1. A phenylene, its hexagonal squeeze and the inner dual of the squeeze.
the hexagonal squeeze of the respective phenylene. An example of a phenylene, its hexagonal squeeze as well as the inner dual of the hexagonal squeeze are shown in Fig. 1.

The graphs described above provide the graph representation of another class of polycyclic hydrocarbons, which are known in chemistry under the name phenylenes. (Using the same name for both a chemical compound and the respective molecular graph should cause no confusion whatsoever.) Phenylenes are very attractive from the point of view of theoretical chemistry because they consist of six-membered cycles (known to stabilize the molecule) and four-membered cycles (known to destabilize the molecule). The (experimental) chemistry of phenylenes is nowadays in rapid expansion (see [32]). It should be mentioned that the number of phenylenes (hydrocarbons) known at present is around 10 , much below the number of known benzenoids. Of the theoretical results on phenylenes (which are not numerous either) we mention the following [12]: If $P H$ is a phenylene and $H S$ is its hexagonal squeeze, then the determinant of the adjacency matrix of $P H$ is equal to (plus or minus) the square of the number of perfect matchings of $H S$. No result on phenylenes (graphs) seems to have ever been reported in the mathematical literature.

## 4. The Wiener number of phenylenes

Let $G$ be a connected graph. Then the relation $\Theta$ on $E(G)$, which was introduced by Djoković [7] in a somewhat different form (see also [10, 20, 21, 35]), is defined as follows. If $e=x y \in E(G)$ and $f=u v \in E(G)$, then $e \Theta f$ if $d(x, u)+d(y, v) \neq d(x, v)+$ $d(y, u)$. The relation $\Theta$ is reflexive and symmetric, yet it need not be transitive. We denote its transitive closure by $\Theta^{*}$ and call the equivalence classes of $\Theta^{*}$ the cuts of $G$. Hence, cuts of $G$ are pairwise disjoint subsets of the edge set $E(G)$.

In Fig. 1 two cuts of the hexagonal squeeze are indicated, as well as the corresponding cuts of the phenylene. In addition to this, one more cut of the phenylene is marked. This cut consists of two opposite (outer) edges of a square. Such cuts will be referred to as square cuts. It is well known that if we remove the edges of a cut from a partial binary Hamming graph (in particular from a hexagonal system or from a phenylene) then the remaining graph consists of two connected components.

Winkler [35] proved that a connected graph $G$ is a partial binary Hamming graph if and only if $G$ is bipartite and $\Theta^{*}=\Theta$. Since hexagonal systems are partial binary Hamming graphs ([23, Theorem 2.1]), it follows that cuts of hexagonal systems coincide with the so-called (elementary, orthogonal) cuts of hexagonal systems, cf. $[23,31,38]$. The same holds for the cuts of phenylenes since as a part of the proof of Theorem 3 we show that phenylenes are partial binary Hamming graphs as well.

Let $H S$ denote the hexagonal squeeze of a phenylene $P H$. Define the weighted graph $\widetilde{H S}=(H S, w)$ by setting $w(u)=2$, if $u$ is a vertex of degree 3 , and $w(u)=1$, otherwise. We say that $\widetilde{H S}$ is the weighted hexagonal squeeze of the phenylene $P H$. For our purpose we can view $\widetilde{H S}$ as a graph which we obtain from $H S$ by replacing each vertex of degree 3 by a pair of vertices at distance 0 , cf. Fig. 2.

We are now prepared to state the following result.


Fig. 2. Another way of representing $\widetilde{H S}$.

Theorem 3. Let PH be a phenylene, let $\widetilde{H S}$ be its weighted hexagonal squeeze and let ID be the inner dual of the hexagonal squeeze. Then

$$
W(P H)=W(\widetilde{H S})+36 W(I D) .
$$

Proof. Let $|V(P H)|=n$ and let $\left\{C_{1}, C_{2}, \ldots, C_{s}, C_{s+1}, C_{s+2}, \ldots, C_{s+t}\right\}$ be the set of cuts of $P H$, where the cuts $C_{i}, s+1 \leqslant i \leqslant t$, correspond to the square cuts and the cuts $C_{i}, 1 \leqslant i \leqslant s$, to the other cuts. Note that $t$ is equal to the number of edges of $I D$ and $s$ is the number of cuts in $H S$.

For $i=1,2, \ldots, s+t$, let $G_{i}^{0}$ and $G_{i}^{1}$ be the connected components of $P H \backslash C_{i}$. Define the $i$ th component of the mapping $g: V(P H) \rightarrow\{0,1\}^{s+t}$ as follows:

$$
g_{i}(u)= \begin{cases}0, & u \in G_{i}^{0} \\ 1, & u \in G_{i}^{1}\end{cases}
$$

Hence, $g(u)=\left(g_{1}(u), g_{2}(u), \ldots, g_{s}(u), g_{s+1}(u), \ldots, g_{s+1}(u)\right)$. Let $u$ and $v$ be any two vertices of $P H$ and let $Q$ be a shortest path in $P H$ between $u$ and $v$. Clearly, if $x y \in C_{\text {, }}$ is an edge of $Q$, then $g_{j}(x) \neq g_{j}(y)$ and this is the only coordinate in which $g(x)$ and $g(y)$ differ. In addition, no two edges of $Q$ belong to the same cut, for otherwise $Q$ would not be a shortest path. It follows that the distance between $u$ and $v$ is equal to the number of coordinates in which $g(u)$ and $g(v)$ differ. Thus $g$ is a binary Hamming labeling of a partial binary Hamming graph $P H$. Then by Corollary 2

$$
W(P H)=\sum_{i=1}^{s+t} n_{i}\left(n-n_{i}\right),
$$

where $n_{i}$ is the number of vertices $u$ of $P H$ with $g_{i}(u)=1$. This equation can be rewritten as

$$
\begin{equation*}
W(P H)=\sum_{i=1}^{s} n_{i}\left(n-n_{i}\right)+\sum_{i=s+1}^{s+t} n_{i}\left(n-n_{i}\right) . \tag{1}
\end{equation*}
$$

Consider the hexagonal squeeze $H S$ of $P H$ and let $C_{1}^{\prime}, C_{2}^{\prime}, \ldots, C_{s}^{\prime}$ be the cuts of $H S$ corresponding to the cuts $C_{1}, C_{2}, \ldots, C_{s}$ of $P H$. Let $H_{i}^{0}$ and $H_{i}^{1}$ be the connected components of $H S \backslash C_{i}^{\prime}$. By [23, Theorem 2.1] $H S$ is a partial binary Hamming graph. Moreover, a binary Hamming labeling $g^{\prime}: V(H S) \rightarrow\{0,1\}^{s}$ can be defined analogously as the labeling $g$ for $P H$, as defined above. Let $u$ be a vertex of $P H$ and let $u^{\prime}$ be the corresponding vertex of $H S$. Then by the definitions of $g$ and $g^{\prime}$ we get that $g^{\prime}\left(u^{\prime}\right)$ is equal to the first $s$ coordinates of $g(u)$. Also, a vertex of $H S$ of degree 3 corresponds to two vertices in $P H$ with identical first $s$ coordinates. By Proposition 1 we hence get

$$
\begin{equation*}
\sum_{i=1}^{s} n_{i}\left(n-n_{i}\right)=W(\tilde{H S}) \tag{2}
\end{equation*}
$$

Consider now the inner dual $I D$ of $H S$. Recall that $I D$ has $t$ edges, say $e_{s+1}, e_{s+2}, \ldots$, $e_{s+t}$, where for $i=s+1, s+2, \ldots, s+t$, the edge $e_{i}$ corresponds to the cut $C_{i}$ of $P H$.

By a classical result of Wiener [33] (or by Corollary 2) we have

$$
W(I D)=\sum_{i=s+1}^{s+1} n_{1}\left(e_{i}\right) n_{2}\left(e_{i}\right)
$$

where $n_{1}\left(e_{i}\right)$ denotes the number of vertices closer to one end vertex of $e_{i}$ and $n_{2}\left(e_{i}\right)$ the number of remaining vertices of $I D$. Let $C_{i}$ be a cut of $P H$ where $i \in\{s+1$, $s+2, \ldots, s+t\}$. Then the number of vertices in $G_{i}^{0}$ is equal to $6 n_{1}\left(e_{i}\right)$ and the number of vertices in $G_{i}^{1}$ is $6 n_{2}\left(e_{i}\right)$, or vice versa. Therefore $n_{i}\left(n-n_{i}\right)=36 n_{1}\left(e_{i}\right) n_{2}\left(e_{i}\right)$ and hence

$$
\begin{equation*}
\sum_{i=s+1}^{s+t} n_{i}\left(n-n_{i}\right)=36 \sum_{i=s+1}^{s+t} n_{1}\left(e_{i}\right) n_{2}\left(e_{i}\right)=36 W(I D) \tag{3}
\end{equation*}
$$

Combining (1) with (2) and (3) completes the proof.

To illustrate Theorem 3 consider the phenylene from Fig. 2. Then $t=6$ (the number of square cuts) and $s=15$ (the number of all the other cuts). By Proposition 1 we have

$$
W(\widetilde{H S})=6 \cdot(3 \cdot 39)+3 \cdot(6 \cdot 36)+3 \cdot(9 \cdot 33)+3 \cdot(18 \cdot 24)=3537
$$

Furthermore, $W(I D)=48$ and by Theorem 3 we conclude that $W(P H)=3537+36$. $48=5265$.

## Acknowledgements

The work of one author (S.K.) was supported in part by the Ministry of Science and Technology of Slovenia under the grant J1-7036. The other author (I.G.) thanks the Mathematical Institute in Belgrade for financial support.

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