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Contact mechanics based solution to predict modulus of asphalt materials with high porosities

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ABSTRACT

Asphalt mixtures with high porosities (known as porous asphalt (PA) mixes) are becoming a popular choice among road authorities as it provides better skid resistance while also reducing tire-pavement noises. Towards the design and manufacture of PA mix pavement, the evaluation of the mechanical properties of PA mixes is of great importance. To predict the mechanical properties of PA mixes, micromechanical models have been considered as an effective tool. In most research studies, continuum-based micromechanical models, i.e. the Self-consistent model, the Mori-Tanaka model, etc. are widely used to predict the stiffness of asphalt mixtures. However, the limitation of these models is that they cannot describe the characteristics of individual particles and thus they cannot provide accurate predictions. On the other hand, the discrete-based micromechanical model (DBMM) which simulates a granular material as an assembly of bonded particles seems to be a promising alternative. Limited research studies have focused on studying the utilization and the applicability of this model for asphalt mixes. Therefore, this paper aims to propose a framework to use DBMM and to evaluate its performance in estimating a PA mix's stiffness. Based on the obtained results, both the merits and limitations of this model were highlighted.

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1. Introduction

Due to the advantage of improving wet skid resistance and reducing hydroplaning and "splash and spray" effects, asphalt mixtures with high porosities, known as porous asphalt (PA) mixes, have attracted the attention of many researchers and engineers [1–4]. PA mixes can be considered as environmental-friendly materials since they are capable of reducing pavement-tire interaction induced noises. All these advantages motivate the wide use of PA mixes in some countries such as the Netherlands where PA mixes surface more than 80% of the highways [5].

In order to come up with a proper PA mix design, a good estimation of the mechanical properties of a given mix is often required. Commonly available pavement analysis tools such as 3D-move [6], together with the mechanical properties of the mix, can be used to estimate the lifetime expectancy of the designed pavements under critical loading conditions. In practice, in order to determine a mix's mechanical properties, a series of laboratory tests are often conducted. However, such laboratory tests are usually not only expensive but also time-consuming.

* Corresponding author. E-mail address: k.anupam@tudelft.nl (K. Anupam). In order to reduce experimental work, researchers [7,8] have made efforts to obtain mechanical properties of an asphalt mixture directly from the properties of its constituents. These empirical/semi-empirical relationships have been broadly used for estimating the mechanical properties of asphalt mixtures [9,10]. However, such models contain various parameters that are calibrated on the basis of a large amount of experimental data. The need for calibration limits the applicability of empirical/semi-empirical models as the determined values of the parameters are only applicable to the mixtures they are calibrated for [11].

With the above realization, researchers started to look for theoretical relationships to predict the mechanical properties of asphalt mixtures. In recent decades, micromechanical models which are developed on the basis of basic mechanical theories (such as continuum mechanics) have drawn the attention of many researchers [12–18]. Among different types of micromechanical models, numerical micromechanical models, which are developed based on finite element method (FEM) and/or discrete element method (DEM), have been favored by many researchers [12–15] mainly because these models can describe almost realistic microstructures of asphalt mixtures (i.e. different sizes and shapes of aggregates). However, the description of such complex microstructures requires the development of a huge number of

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finite or discrete elements. This indicates that powerful computational facilities are always needed to analyze the mechanical properties of asphalt mixtures, and it always takes a long time to obtain the predicted results.

Furthermore, although the microstructure of asphalt mixtures can be realistically described using FEM or DEM, accurate predictions still rely on the calibration of some input parameters. For example, experimental results are typically needed to calibrate the parameters of the contact constitutive models between two particles when DEM is used to simulate the mechanical properties of asphalt mixtures [12,13].

On the other hand, analytical micromechanical models can be a promising alternative to determine the mechanical properties of asphalt mixtures in a more effective way. These models simplify the descriptions of the microstructure of asphalt mixtures. For example, aggregates are typically modelled as identical spherical or ellipsoidal particles. However, these simplified descriptions allow obtaining analytical solutions to the mixture's mechanical properties. Therefore, in comparison to FEM and DEM, analytical micromechanical models are much less time and facilitiesconsuming. Furthermore, although analytical micromechanical models cannot realistically describe the microstructure of asphalt mixtures, their capabilities of accurately predicting the mechanical properties of the mixtures have been reported in many research studies [19,20].

Currently, research work regarding analytical micromechanical models has mainly focused on the continuum-based micromechanical models (CBMM), such as the Self-consistent (SC) model [21], the Mori-Tanaka (MT) model [22], the Differential model [23], the generalized Self-consistent (GSC) model [24], etc. These models are developed based upon a matrix-inclusion system where an inclusion is embedded into a continuous matrix [25]. Although CBMM could provide accurate predictions for materials with low concentrations of particles, the predictions at high concentrations did not match well with the experimental results, especially at low frequencies [16–18].

Researchers postulated that the inaccuracies of the predictions could be related to the fact that CBMM do not sufficiently account for the stiffening effect of particles' interactions [16,17]. In a composite, particles' interactions are always associated with the characteristics of individual particles, such as their sizes, shapes, locations, etc. On the contrary in CBMM, all the particles are represented as one phase, which makes it impossible for the models to include these characteristics.

In recent years, few micromechanical models where a specific microstructure of a composite can be defined have also been used by pavement researchers [26]. For example, in the physical interaction model [26], different sizes of individual particles are embedded into a continuous matrix, see Fig. 1. On the basis of the distribution of the particles and their different sizes, the stiffness



Fig. 1. Illustration of the physical interaction model.

of the composite can be estimated by quantifying the stiffening effects of individual particles and their interactions.

Although the physical interaction model provides a way to consider the characteristics of individual particles, it is only suitable for composites without or with few air voids, such as mastic and dense asphalt mixtures. This is because the matrix in the model is considered to be continuous. However, in a PA mix, since there is a high content of air voids, the matrix phase is expected to be discontinuously located among densely packed aggregate particles. Therefore, in order to accurately predict the mix's mechanical properties, a different micromechanical model which can reasonably describe the microstructure of the mix is required.

In comparison to the physical interaction model, the discretebased micromechanical model (DBMM) [27–29] potentially provides a better way to describe the particles' interactions in PA mixes. This model simulates a bonded granular material as a particle-on-particle skeleton bonded by binder layers. This arrangement is naturally expected to represent a more reasonable and realistic microstructure for PA type mixes. Moreover, such a model has the capability to include several geometrical parameters to describe the characteristics of individual particles, for example, the radius of the particles, the average number of contacts per particle, etc.

Solutions to the effective moduli of a bonded granular material using DBMM have been derived by different researchers [27,29]. In this study, solutions provided by Dvorkin, et al [27] were used and thus DBMM will be referred to as Dvorkin's model in the following sections. Dvorkin's model has been successfully used to predict the modulus of granular materials bonded by elastic binders such as frozen sand, glass beads packs, etc. [30]. According to the elasticviscoelastic corresponding principle [31], models for elastic materials can be used for viscoelastic materials as well. Therefore, it is highly possible that Dvorkin's model could be used for estimating the modulus of asphalt materials. However, limited research studies have provided thorough investigations in this area. For example, Zhu and Nodes [28] used Dvorkin's model to simulate the creep behaviour of asphalt mixtures, limiting their study to some randomly assumed geometrical parameters, whereas, it is obvious that the values of geometrical parameters significantly affect the predictions.

In light of the above discussions, the main aim of this study is to investigate the performance of Dvorkin's model on predicting the modulus of PA mixes. The scope of this study includes:

- to propose a framework for predicting the modulus of PA mixes using Dvorkin's model;
- to evaluate the performance of Dvorkin's model based upon the experimental results of the complex modulus of PA mixes.

In order to make this paper understandable for readers, it has been arranged in different sections. The first and second sections will introduce and provide background information about Dvorkin's model. The proposed framework for predicting the effective moduli of PA mixes using Dvorkin's model will be elaborated in the subsequent section. In the fourth section, materials characteristics and performed laboratory tests will be presented. The fifth section will discuss the obtained model parameters. The last two sections will describe the predicted results and conclusions.

2. Background knowledge

2.1. Dvorkin's model

In Dvorkin's model, the overall effective moduli of a granular material are derived by homogenizing the stiffness of an assembly of two-bonded particle systems that orient in different directions. This section first describes the stiffness of a two-bonded particle system, on the basis of which, a homogenization technique is introduced to obtain the effective (homogenized) moduli of a bonded granular material.

2.1.1. Stiffness of a two-bonded particle system

A two-bonded particle system consisting of two equal-sized spheres and a layer of binder is illustrated in Fig. 2a. The spherical particles are assumed to have an arbitrary radius R, with the binder layer radius of a and a minimum height of 2 h_0 . It is noted here that generally, particles in a granular material have different sizes and irregular shapes. Since closed-form solutions for calculating the stiffness of a two-bonded particle system with irregular shapes and different sizes have not yet been accurately derived, past researchers have considered these particles as spheres with uniform sizes [27,29].

In Dvorkin's model, the stiffnesses of a two-bonded particle system *S* are defined as:

$$F_n = S_n \delta_n, F_\tau = S_\tau \delta_\tau \tag{1}$$

where *F* is the applied force on the system; δ is the displacement of the centre of one particle relative to the median plane (the *x* axis); and the subscripts *n* and τ denote the normal and the tangential direction, respectively.

The value of δ is related to the displacement of the binder layer's surface (represented as *V* and *U* in the normal direction and the tangential direction, respectively) as well as that of the sphere's surface (represented as *v* and *u* in the normal direction and the tangential direction, respectively), see Eqs. (2) and (3).

$$\delta_n = v(\mathbf{x}) - V(\mathbf{x}) \tag{2}$$

$$\delta_{\tau} = u(x) - U(x) \tag{3}$$

In order to calculate the displacement of the binder layer, researchers [27] proposed a solution for a similar problem assuming a thin binder layer as an elastic base. In this solution, V(x) and U(x) were related to the stress along the interface σ via Eqs. (4) and (5), respectively. Furthermore, since the contact region is small as compared to the particle, v(x) and u(x) can be approximated as the surface's displacement of an elastic half-space [27], see Eqs. (6) and (7), respectively.

$$\sigma_n(x) = -\frac{2G_b(1-v_b)}{1-2v_b} \frac{V(x)}{h(x)}$$
(4)

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$$\sigma_{\tau}(\mathbf{x}) = -G_{\rm b} \frac{U(\mathbf{x})}{h(\mathbf{x})} \tag{5}$$

$$\nu(x) = \frac{1 - \nu_p}{\pi G_p} \int_0^{\pi} d\varphi \int_0^{\beta} \sigma_n(r) ds \tag{6}$$

$$u(x) = \frac{1}{2\pi G_{\rm p}} \int_0^{\pi} d\phi \int_0^{\beta} \sigma_{\tau}(r) (1 - v_{\rm p} \sin^2 \phi) ds$$
(7)

where

$$\beta = x\cos\varphi + \sqrt{a^2 - x^2\sin^2\varphi}$$

$$r = \sqrt{x^2 + s^2 - 2xs\cos\varphi}$$
vs and Gs are the Poisson's ratio and the shear modulus of th

 v_b and G_b are the Poisson's ratio and the shear modulus of the binder layer, respectively; v_p and G_p are the Poisson's ratio and the shear modulus of the particles, respectively; h(x) is the half-thickness of the binder layer, which can be approximated as

$$h(x) = h_0 + \frac{x^2}{2R}$$
(8)

By combining Eqs. (2), (4) and (6) or Eqs. (3), (5) and (7), the displacement of the binder layer's surface can be obtained giving a non-zero constant value of δ , see Eqs. (9) and (10). These two equations are known as the Volterra integral equations of the second kind. Since this type of integral equations is too complicated to be solved analytically, a numerical technique was used, see Appendix A. Once the displacement of the binder layer's surface is known, the value of σ can be computed from Eqs. (4) and (5). The value of *F* is determined by integrating σ on the whole area of the interface, see Eqs. (11) and (12).

$$\delta_n + V(x) = \Lambda_n \int_0^\pi d\varphi \int_0^\beta \frac{V(r)}{h(r)} ds$$
(9)

$$\delta_{\tau} + U(x) = \Lambda_{\tau} \int_0^{\pi} d\varphi \int_0^{\beta} \frac{U(r)}{h(r)} (1 - v_{\rm p} \sin^2 \varphi) ds \tag{10}$$

where

$$\Lambda_n = -\frac{2G_{\rm b}}{\pi G_{\rm p}} \frac{(1-\nu_{\rm p})(1-\nu_{\rm b})}{1-2\nu_{\rm b}}$$

$$\Lambda_{ au} = -rac{G_b}{2\pi G_p}$$



a. Two bonded particles

b. Stress distribution at the interface

Fig. 2. A two-bonded particle system.

$$F_n = \int_0^a \sigma_n(\mathbf{x}) \cdot 2\pi \mathbf{x} \cdot d\mathbf{x} \tag{11}$$

$$F_{\tau} = \int_{0}^{a} \sigma_{\tau}(x) \cdot 2\pi x \cdot dx \tag{12}$$

It is noted that the above solutions were developed for particles bonded by an elastic binder layer. According to the elastic–viscoelastic corresponding principle, these solutions can be used for viscoelastic materials by replacing the elastic moduli of the binder layer with the complex moduli of the viscoelastic binder layer.

2.1.2. Homogenization technique

In order to upscale the mechanical properties of a two-bonded particle system to the effective moduli of a granular material, a homogenization technique is needed. As the first step, an assembly of bonded particles, as illustrated in Fig. 3, is considered as a representative volume element (RVE) of a bonded granular material. In the RVE, the particles are considered as spheres with a uniform radius of R; and all the binder layers are assumed to have a radius of a and a minimum height of 2 h_0 .

Two spheres *r* and *s*, which are initially in contact with each other, have the position vectors of $\mathbf{X}^{(r)}$ and $\mathbf{X}^{(s)}$ in the global coordinate system, respectively. Under a deformation **u** subjected to the boundary of the system, the centers of *r* and *s* also undergo displacements, represented as $\mathbf{u}^{(r)}$ and $\mathbf{u}^{(s)}$, respectively. By using the kinematic hypothesis that the strains throughout the packing are uniform, the values of $\mathbf{u}^{(r)}$ and $\mathbf{u}^{(s)}$ can be calculated via Eq. (13).

$$u_i^{(r)} = \left\langle \epsilon_{ij} \right\rangle : X_j^{(r)}, u_i^{(s)} = \left\langle \epsilon_{ij} \right\rangle : X_j^{(s)}$$
(13)

where $\langle \varepsilon_{ij} \rangle$ is the average strain applied on the packing.

By symmetry, it can be obtained that the center of the median plane between the *s*th and the *r*th particles undergoes a displacement of $(\mathbf{u}^{(r)} + \mathbf{u}^{(s)})/2$. Thus, relative to the median plane's center point, the displacement of the *r*th sphere center is $(\mathbf{u}^{(r)}-\mathbf{u}^{(s)})/2$. This relative displacement can be separated into a normal component δ_n (Eq. (14)) and a shear component δ_{τ} (Eq. (15)). The total force **F** on the *s*th sphere due to its contact with the *r*th sphere is given as the sum of the force in the normal direction and that in the tangential direction, see Eq. (16). By substituting Eq. (13) into Eq. (16), the value of **F** can be further written as Eq. (19).

$$\boldsymbol{\delta}_{n} = \left[\frac{1}{2} \left(\boldsymbol{u}^{(r)} - \boldsymbol{u}^{(s)}\right) \cdot \boldsymbol{I}^{(sr)}\right] \boldsymbol{I}^{(sr)}$$
(14)



$$\boldsymbol{\delta}_{\tau} = \frac{1}{2} \left(\boldsymbol{u}^{(r)} - \boldsymbol{u}^{(s)} \right) - \boldsymbol{\delta}_{n}$$
(15)

$$\boldsymbol{F}^{(sr)} = (\boldsymbol{S}_n - \boldsymbol{S}_{\tau}) \left[\frac{1}{2} \left(\boldsymbol{u}^{(r)} - \boldsymbol{u}^{(s)} \right) \cdot \boldsymbol{I}^{(sr)} \right] \boldsymbol{I}^{(sr)} + \boldsymbol{S}_{\tau} \left[\frac{1}{2} \left(\boldsymbol{u}^{(r)} - \boldsymbol{u}^{(s)} \right) \right]$$
(16)

$$\boldsymbol{F}^{(sr)} = (S_n - S_\tau) \left[\frac{1}{2} \langle \boldsymbol{\varepsilon} \rangle : \left(\boldsymbol{X}^{(r)} - \boldsymbol{X}^{(s)} \right) \cdot \boldsymbol{I}^{(sr)} \right] \boldsymbol{I}^{(sr)} + S_\tau \left[\frac{1}{2} \langle \boldsymbol{\varepsilon} \rangle : \left(\boldsymbol{X}^{(r)} - \boldsymbol{X}^{(s)} \right) \right]$$
(17)

Furthermore, it is defined that **I** is the unit vector along the line of centers of two particles:

$$\mathbf{I}^{(sr)} = \frac{\mathbf{X}^{(s)} - \mathbf{X}^{(r)}}{2(R+h_0)}$$
(18)

Substituting Eq. (18) into Eq. (17), the value of **F** can be further given as

$$F_{i}^{(\mathrm{sr})} = (S_{n} - S_{\tau}) \left(-(R + h_{0}) \langle \varepsilon \rangle_{kl} I_{k}^{(\mathrm{sr})} I_{l}^{(\mathrm{sr})} \right) I_{i}^{(\mathrm{sr})} + S_{\tau} \left(-(R + h_{0}) \langle \varepsilon \rangle_{ij} I_{j}^{(\mathrm{sr})} \right)$$
(19)

The average stress field $\langle \sigma \rangle$ of the granular material with a total volume *V* is related to the stresses within individual particles by using the following equation:

$$\langle \sigma \rangle_{ij} = \frac{1}{V} \sum_{N_p} \int_{V_s} \sigma_{ij}^{(s)} dV \tag{20}$$

where $\sigma_{ij}^{(s)}$ is the stress within the *s*th particle; *V_s* represents the volume of the sphere and *N_p* is the number of all the particles within *V*.

By using the divergence theorem, the value of $\sigma_{ij}^{(s)}$ can be calculated from the traction on the surface of the particle:

$$\int_{V_s} \sigma_{ij}^{(s)} dV = \frac{1}{2} \int_{S_s} \left(x_i' \tau_j^{(s)} + x_j' \tau_i^{(s)} \right) dS$$
(21)

where S_s is the surface of the s^{th} particle; **x**' denotes the position vector of a point on S_s relative to the center of the particle; $\tau^{(s)}$ denotes the traction across S_s . From Fig. 3, it can be seen that the value of $\tau^{(s)}$ is nonzero only at the position where the particle contacts with other particles. Therefore, Eq. (21) can be further written as

$$\int_{V_s} \sigma_{ij}^{(s)} dV = \frac{1}{2} \sum_n \left(x'_i F_j^{(rs)} + x'_j F_i^{(rs)} \right)$$
(22)

where n is the number of contacts per particle.

Since the contact area between the s^{th} and the r^{th} particle is small, the value of **x'** at the contact area can be approximated by the position vector of the center of the contact area relative to the center of the s^{th} particle:

$$\int_{V_s} \sigma_{ij}^{(s)} dV = \frac{1}{2} \sum_n \left[\frac{1}{2} \left(X_i^{(r)} - X_i^{(s)} \right) F_j^{(rs)} + \frac{1}{2} \left(X_j^{(r)} - X_j^{(s)} \right) F_i^{(rs)} \right]$$
(23)

By substituting Eq. (23) into Eq. (20), the value of $\langle\sigma\rangle$ can be written as

$$\langle \sigma \rangle_{ij} = \frac{1}{2V} \sum_{N_p} \sum_{n} \left[\frac{1}{2} \left(X_i^{(r)} - X_i^{(s)} \right) F_j^{(rs)} + \frac{1}{2} \left(X_j^{(r)} - X_j^{(s)} \right) F_i^{(rs)} \right]$$
(24)

This value can be further expressed as Eq. (25) by substituting Eq. (19) and Eq. (18) into Eq. (24).

$$\langle \sigma \rangle_{ij} = \frac{\left(R + h_0\right)^2}{2V} \sum_{N_p} \sum_{n} \left[\frac{2(S_n - S_\tau) \left(\langle \varepsilon \rangle_{kl} I_k^{(sr)} I_l^{(sr)} \right) I_i^{(rs)} I_j^{(rs)}}{+S_\tau \left(\langle \varepsilon \rangle_{jk} I_k^{(rs)} I_i^{(rs)} + \langle \varepsilon \rangle_{ik} I_k^{(rs)} I_j^{(rs)} \right)} \right]$$
(25)

Assuming that the geometry of the packing is statistically isotropic and the distribution probability of the contact points over the surface of a sphere is uniform, the summation in Eq. (25) can be represented in terms of averages:

$$\left\langle \sigma \right\rangle_{ij} = \frac{\left(R + h_0\right)^2 n N_p}{2V} \left[2(S_n - S_\tau) \left\langle \langle \varepsilon \rangle_{kl} I_k I_l I_l J_j \right\rangle + S_\tau \left(\left\langle \langle \varepsilon \rangle_{jk} I_k I_j \right\rangle + \left\langle \langle \varepsilon \rangle_{ik} I_k I_j \right\rangle \right) \right]$$
(26)

where the brackets <-> denote the average over all uniformly distributed unit vector **I**.

In Eq. (26), the total volume of the system V can be given as

$$V = \frac{V_{\rm p}}{\phi_{\rm p}} \tag{27}$$

where V_p and ϕ_p denote the volume and the volume fraction of the particles in the RVE, respectively. When all the particles are spheres with uniform sizes, the value of V_p is given as

$$V_{\rm p} = \frac{4}{3}\pi R^3 N_{\rm p} \tag{28}$$

Combining Eqs. (27) and (28), the value of $N_{\rm p}/V$ can be expressed as

$$\frac{N_{\rm p}}{V} = \frac{3}{4} \frac{\phi_{\rm p}}{\pi R^3} \tag{29}$$

Therefore, Eq. (26) can be further written as

$$\langle \sigma \rangle_{ij} = \frac{3n\phi_{p}(R+h_{0})^{2}}{8\pi R^{3}} \left[2(S_{n}-S_{\tau}) \left\langle \langle \varepsilon \rangle_{kl} I_{k} I_{l} I_{l} I_{j} \right\rangle + S_{\tau} \left(\left\langle \varepsilon \rangle_{jk} \left\langle I_{k} I_{i} \right\rangle + \left\langle \left\langle \varepsilon \right\rangle_{ik} I_{k} I_{j} \right\rangle \right) \right]$$
(30)

2.1.3. Effective moduli of a bonded granular material

The stiffness tensor C^* of a bonded granular material can be derived from Eq. (30) as

$$C_{ijkl}^{*} = \frac{3n\phi_{p}(R+h_{0})^{2}}{16\pi R^{3}} \left[4(S_{n}-S_{\tau})\langle I_{i}I_{j}I_{k}I_{l}\rangle + S_{\tau} \left(\langle I_{k}I_{i}\rangle\delta_{jl} + \langle I_{k}I_{j}\rangle\delta_{il} + \langle I_{l}I_{i}\rangle\delta_{jk} + \langle I_{l}I_{j}\rangle\delta_{ik} \right) \right]$$
(31)

According to the relationships in Eq. (32) and the definition of the stiffness tensor for an isotropic material in Eq. (33), the Young's modulus E^* of a bonded granular material can be calculated using Eq. (34). By substituting the determined values of S_n and S_τ from Eq. (1) into Eq. (34), the value of E^* can be obtained.

$$\langle I_i I_j \rangle = \frac{1}{3} \delta_{ij}, \langle I_i I_j I_k I_l \rangle = \frac{1}{15} \left(\delta_{ij} \delta_{kl} + \delta_{ik} \delta_{jl} + \delta_{ik} \delta_{jk} \right)$$
(32)

$$C_{ijkl}^{*} = \frac{G^{*}(2G^{*} + E^{*})}{3G^{*} - E^{*}} \delta_{ij}\delta_{kl} + G^{*} \left(\delta_{ik}\delta_{jl} + \delta_{il}\delta_{jk}\right)$$
(33)

$$E^* = \frac{n\phi_{\rm p}(R+h_0)^2}{4\pi R^3} \frac{2S_n + 3S_{\tau}}{4S_n + S_{\tau}} S_n$$
(34)

where *n* denotes the coordination number; ϕ_p is the volume fraction of the particles; *R* represents the radius of the particles; h_0 refers to the minimum distance between adjacent particles; and S_n and S_τ denote the stiffnesses of a two-bonded particle system in the normal and in the tangential direction, respectively.

3. Proposed framework to predict stiffness of PA mixes

The proposed framework for predicting the stiffness of PA mixes contains three main steps, see Fig. 4. A PA microstructure was assumed to be consisting of randomly packed spherical particles, mortar, and air voids. The total volume of mortar was categorized into two functional groups. Some mortar layers that are located between adjacent particles play a major role in binding particles

together, and thus they were defined as the "binding mortar layers"; whereas the remaining parts only play a role in coating individual particles, and thus they were defined as the "coating mortar layers".

To predict the stiffness of PA mixes, the mechanical, volumetric and geometric properties of every phase are required. Generally, the mechanical and the volumetric properties can be directly measured from laboratory tests. Additionally, five geometric parameters need to be determined: (1) the radius of the spherical particles, (2) the thickness of the coating mortar layers, (3) the minimum thickness of the binding mortar layers, (4) the average coordination number, and (5) the radius of the binding mortar layers. These geometric parameters may also be measured using advanced technologies such as digital image processing. However, this method is difficult to be implemented since the values of all these geometric parameters vary with different locations. Thus, in this study, a different method that can be used to determine the geometric parameters of the proposed microstructure in a much easier but reasonable way was proposed, see Appendix B.

In the proposed microstructure, the properties of both the binding mortar layers and the coating mortar layers affect the stiffness of a PA mix. However, Dvorkin's model can only predict the stiffness of a skeleton framework consisting of particles, air voids and the binding mortar layers. Therefore, in the last step, a different methodology was employed to include the effect of the coating mortar layers in the model.

3.1. Proposed microstructure model for PA mixes

The microstructure model for PA mixes, see Fig. 5, was proposed according to the literature [32]. Individual particles (modelled as identical spheres with a uniform radius of R) are initially covered by mortar layers with a uniform thickness of t. When the mortar-covered aggregate particles pack together, the minimum distance between adjacent particles, represented as 2 h_0 , has an initial value of 2 t, see Fig. 5a. It was assumed that during the compaction process, the value of h_0 decreases, and depending on the amount of the compaction work, the value of h_0 ranges from t (without any compaction effort) to 0 (two particles contact each other), see Fig. 5b. It is noted here that during the compaction, not only the value of h_0 decreases but also the relative positions of aggregate particles change. However, since the change of the particles' positions is difficult to be analytically described, the proposed microstructure model did not take this phenomenon into account.

The volume of the binding mortar layer between two adjacent particles is illustrated in Fig. 6a. The binding mortar layers, with a radius of a, are assumed to be composed of two parts, see Fig. 6b. When two particles are compacted closer, their coating mortar layers overlap each other to form one part of the binding mortar layer with a radius of a_1 . Due to the overlap of the coating mortar layers, some mortar is squeezed out and forms the other part of the binding mortar layer with a radius of a_2 .

3.2. Procedures to calculate the stiffness of PA mixes

According to the literature [30], contributions to the stiffness of a mix made by the binding mortar layers and the coating mortar layers are different. The binding mortar layers play a major role in providing the stiffness of the mix because they enable the discrete particles to work together as a skeleton framework. By contrast, the coating mortar layers contribute to the stiffness of the mix by filling the air voids in the skeleton framework.

In light of the above discussions, the following procedure was proposed to calculate the stiffness of PA mixes. At first, the stiffness of the skeleton framework consisting of aggregate particles and the



Fig. 4. Flowchart of the proposed framework.



Fig. 5. Proposed microstructure model for PA mixes.



Fig. 6. Illustration of binding mortar layers.

binding mortar layers was predicted using Dvorkin's model, see Fig. 7a. This skeleton framework was further equivalent as a twophase composite with air voids embedded into a continuous matrix. This two-phase composite's stiffness was assumed to be the same as the skeleton framework. The air voids' volume fraction in the two-phase composite was equal to the sum of the volume fractions of the air voids and the coating mortar layers in the PA mix. Based on the stiffness of the composite and the volume fraction of each phase, the matrix's stiffness can be back-calculated.

At last, the coating mortar layers were added into the skeleton framework to form the PA mix. This was equivalent to the process that the same volume of mortar was embedded into the matrix of the two-phase composite. As a result, a three-phase composite, consisting of the matrix, coating mortar layers and air voids, was formed, see Fig. 7b. The volume fractions of the coating mortar layers and the air voids in this three-phase composite were the same as those in the PA mix. Based on the stiffness and the volume fraction of each phase, the stiffness of the three-phase composite was calculated. The stiffness of the PA mix, which was equal to the stiffness of the three-phase composite, became known as well.

Since the matrix in the two-phase composite and the threephase composite was continuous, the back-calculation of the matrix's stiffness and the calculation of the stiffness of the threephase composite can be achieved using CBMM. In this study, the SC model was preferred over others due to the following reasons: (a) this model can be easily solved to back-calculate the stiffness of the matrix (comparing to the GSC model and the Differential model) [33]; and (b) it can provide more accurate predictions when the volume fraction of inclusions is lower than 50% (comparing to the Dilute model and the MT model) [34]. Appendix B describes the procedure to back-calculate the matrix's stiffness of a two-phase composite using the SC model.

Up to now, previous sections briefly introduced Dvorkin's model and presented a framework to predict the stiffness of PA mixes. The next part of the research was to evaluate the performance of Dvorkin's model on the basis of laboratory investigation. Using a Dynamic Shear Rheometer (DSR) device, frequency sweep tests were conducted on laboratory prepared mortar specimens to obtain input parameter $G_{\rm b}$; while the PA mix specimens' complex Young's modulus $E_{\rm mix}$, which was used to evaluate the performance of the model, was measured from uniaxial cyclic compressive tests.

4. Materials and tests

4.1. Specimen preparation

4.1.1. PA mix specimens

In this study, in order to evaluate the performance of Dvorkin's model for PA mixes with different skeleton frameworks, two types of PA mix materials (PA mix-A and PA mix-B) were considered. PA mixes were designed according to the Dutch standard [35,36]. For both mixes, the gradation of the aggregates and their respective densities are shown in Table 1, and the content of asphalt binder with a penetration of 70–100 was 4.3%. The density of the binder was assumed as 1030 kg/m³.

Three replicates of PA mix specimens for uniaxial compressive tests were prepared following the AASHTO T 342-11 standard method [37]. A gyratory compactor was firstly used to make PA mix specimens with a size of Φ 150 mm × H170 mm. In order to generate different skeleton frameworks, different compaction efforts were performed. The number of the gyrations for making PA mix-A specimens was 5 cycles while for PA mix-B, 40 cycles were performed. As a result, a more air voids content of 18% was achieved for PA mix-A whereas the air voids content of PA mix-B was 13.2%. Since the final size of PA mix specimens for stiffness measurement is 100 mm in diameter and 150 mm in height, specimens from the gyratory compactor were further cored and cut.



a. Skeleton framework equivalent as a two-phase composite



b. PA mix equivalent as a three-phase composite

Fig. 7. Procedures to calculate the stiffness of PA mixes.

Table 1

Gradation of aggregates in the PA mix specimens.

Size (mm)	22.4	16	11.2	8	5.6	2	0.5	0.063	filler
Gradation (% Passing)	100	98	77	44	22	15	14	4	0
Density (kg/m ³)	2686	2686	2686	2678	2670	2673	2658	2658	2638

4.1.2. Mortar specimens

All the volumes of the asphalt binder, filler and sand (<0.5 mm) in the mix were used to make mortar specimens. By normalizing the proportion of each size of the sand with respect to the maximum size, the gradation of the sand in the mortar was obtained, see Table 2. The asphalt binder's content in the mortar was computed as 23% according to the content of asphalt binder (4.3%) and the proportion of the sand ((1–4.3%)×14% = 13.4%) in the mix.

The dimension of the mortar specimens is shown in Fig. 8a. The middle part, with a diameter of 6 mm and a height of 12 mm, is used to measure the material's properties; while the end parts, where steel rings with a height of 4 mm and a thickness of 1 mm were fixed, are used to clamp the specimens on the testing machine. A special mould was used to make these specimens, see Fig. 8b. It is highlighted that due to the high asphalt binder content, when the mortar material was heated up to around 160 °C, it could smoothly flow. Therefore, there was no compaction work applied during the preparation. More details on the preparation of mortar specimens can be found in reference [38].

4.2. Laboratory tests

4.2.1. Frequency sweep test

Frequency sweep tests with a frequency range from 20 Hz to 0.1 Hz were conducted at five different temperatures of -10 °C, 4 °C, 21 °C, 37 °C and 54 °C. At each temperature, in order to guarantee the mortar behaves as a linear viscoelastic material, a constant small strain from 10µε to 200µε was used.

4.2.2. Uniaxial cyclic compressive test

Universal Testing Machine (UTM), see Fig. 9, was used to conduct uniaxial cyclic compressive tests at seven different temperatures, -10 °C, 4 °C, 21 °C, 37 °C, 45 °C, 54 °C and 60 °C, and six different frequencies of 20 Hz, 10 Hz, 5 Hz, 1 Hz, 0.5 Hz and 0.1 Hz. Forces were measured via the load cell at the top of the specimen and the displacement was measured via three Linear Variable Differential Transformer sensors. During all the tests, a constant strain with an amplitude of $10\mu\epsilon$ was controlled.

5. Specified values of the input parameters in Dvorkin's model

Using the proposed method in Section 3, the values of the parameters (i.e. mechanical parameters, volumetric and geometric parameters) in the proposed microstructure were determined on the basis of the materials' properties. In this section, these determined values will be presented.

5.1. Mechanical parameters

The master curves of the dynamic shear modulus $|G_b|$ and the phase angle δ_b of the mortar at a reference temperature of 21 °C were constructed according to the time-temperature

superposition principle, see Fig. 10. It can be observed that with the decrease of frequencies, the value of $|G_b|$ keeps decreasing to be close to zero, and meanwhile the value of δ_b increases to be approximately 90°. This can be related to the fact that the volume fraction of the fine particles in mortar is not high enough to form a particle-on-particle skeleton, and thus its behavior is governed by asphalt binder.

The values of other mechanical parameters v_b , G_p and v_p were obtained from the literature [16,39] as $v_b = 0.4$, $G_p = 20.8$ GPa and $v_{agg} = 0.27$. It is highlighted that the value of v_b is expected to be sensitive to test frequency and temperature. However, due to the difficulty of measuring the radial deformation of a mortar specimen, it was difficult to directly measure v_b from laboratory tests. Therefore, a constant value of v_b between 0.35 and 0.5 is generally assumed in the literature [16].

5.2. Volumetric and geometrical parameters

As mentioned earlier, the values of ϕ_v for PA mix-A and PA mix-B were 18% and 13.2%, respectively. According to the content and the density of asphalt binder, and the aggregate gradation and their respective densities, the values of ϕ_b and ϕ_p for PA mix-A were calculated as 20% and 62%, respectively, while for PA mix-B, these values were 20.8% and 66%, respectively.

Since PA mix-A and PA mix-B contain the same aggregate gradations and the same asphalt binder contents, the values of *R* and *t* are identical for both mixes. The value of *R* was calculated as 4 mm using Eq. (B.1). The value of β was calculated as 0.24 from the values of $\phi_{\rm b}$ and $\phi_{\rm p}$ using Eq. (B.3). Once the value of β was known, the value of *t* in Eq. (B.5) was computed as 0.39 mm.

As can be seen in Fig. B.2, when the air voids contents are 18% and 13.2%, the values of h_0/t are approximately equal to 0.58 and 0, respectively. Therefore, the values of h_0 were estimated as 0.22 mm and 0 mm for PA mix-A and PA mix-B, respectively. Using the values of $\phi_{\rm v}$, R, t and $h_{\rm 0}$, the values of $\phi_{\rm 0}$ were calculated as 0.268 and 0.340 using Eq. (B.12) for PA mix-A and PA mix-B, respectively. The values of n were further obtained as 11.5 and 9.5 for PA mix-A and PA mix-B, respectively. On the basis of the obtained values of R, t and h_0 , the values of a were computed as 1.67 mm and 2.53 mm using Eq. (B.16) for PA mix-A and PA mix-B, respectively. Furthermore, using the values of $\phi_{\rm b}$, R, t, h_0 , n and a, the values of $\phi_{\rm b_b}$ were computed from Eqs. (B.18)-(B.19) as 9.4% and 20.3% for PA mix-A and PA mix-B, respectively. The values of $\phi_{\rm b_c}$ were further obtained using Eq. (B.20) as 10.6% and 0.7% for PA mix-A and PA mix-B, respectively. The calculated results of all the parameters as well as the number of the figures and/or equations that were used for determining these parameters are summarized in Table 3.

Comparing the values of the geometric parameters between PA mix-A and PA mix-B, it can be found that different compaction effort affects individual phases in terms of not only their volumetric properties in the macroscale but also their geometric properties

Tubic				
Sand	gradation	in	the	mortar.

Table 2

Size (mm)	0.5	0.18	0.125	0.063	Filler
Gradation (% Passing)	100	62	39	29	0



a. Mortar specimens' dimension



b. Mould for making mortar specimens

Fig. 8. Mortar specimens' preparation.



Fig. 9. Uniaxial cyclic compressive tests for PA mixes.

in the microscale. A larger amount of compaction effort decreases the average distance between adjacent particles (or the average thickness of the binding mortar layers). When two particles become closer, a higher proportion of the coating mortar layers overlap each other to form the binding mortar layers. Therefore,

Table 3	1					
Values	of volumetric	and	geome	tric param	eters.	

Parameters	PA mix-A	PA mix-B	Equations and/or Figures
ϕ_{b}	20%	20.8%	NA
$\phi_{\rm p}$	62%	66%	NA
ϕ_{v}	18%	13.2%	NA
R	4 mm	4 mm	Eq. (B.1)
t	0.39 mm	0.39 mm	Eq. (B.5)
ho	0.22 mm	0 mm	Fig. B.2
n	11.5	9.5	Eqs. (B.9) and (B.12)
а	1.67 mm	2.53 mm	Eq. (B.16)
ϕ_{b_b}	9.4%	20.3%	Eqs. (B.17)–(B.19)
$\phi_{b_{-c}}$	10.6%	0.7%	Eq. (B.20)

with the decrease of the distance between adjacent particles, the radius, as well as the total volume of the binding mortar layers, increases. On the other hand, more compaction effort decreases the average coordination number. This can be explained by the fact that when the particles become closer, less space around one particle is provided for other particles to surround it.

6. Results and discussions

6.1. Predicted results of PA mixes' modulus

In Fig. 11, both the experimental results and the predicted values of E_{mix} (represented in terms of dynamic Young's modulus | E_{mix} | and phase angle δ_{mix}) are presented. It can be seen that the experimental values of $|E_{\text{mix}}|$ show asymptotic behaviour both at very high frequencies and very low frequencies. In addition, with the decrease of frequencies, the experimental results of δ_{mix}



Fig. 10. Measured shear modulus of mortar.



Fig. 11. Comparison of tested and predicted E_{mix.}

increase initially and decrease after a peak point. It is highlighted here that previous research work [38], concluded that these behaviours of PA mixes are strongly related to the characteristics of the stone-on-stone skeleton in the mix.

Fig. 11 also shows a comparison between the experimental and the predicted values. From the plots, it can be seen that Dvorkin's model can predict E_{mix} and δ_{mix} quite well for frequencies higher than around 0.1 Hz. This shows that Dvorkin's model works well within specified frequencies.

In order to check the viability of Dvorkin's model, predicted results were compared against commonly used CBMM in Fig. 11. The Differential model was chosen over other models because previous research studies showed that the Differential model provided better predictions for PA mixes [17]. As can be seen in Fig. 11, at higher frequencies, for a loosely compacted PA mix (i.e. PA mix-A), the predicted results of $|E_{mix}|$ and δ_{mix} do not show significant differences between Dvorkin's models and the Differential model. Whereas, for a densely compacted PA mix (PA mix-B), Dvorkin's model provides better predictions. This can be related to the fact that Dvorkin's model is developed mainly for densely packed granular materials [30]. Apart from the volumetric properties of each phase, Dvorkin's model also takes into account the stiffening effects of the geometric characteristics of individual particles and mortar layers. Therefore, in comparison to the Differential model, Dvorkin's model can account for the inter-particle interactions more accurately.

The limitation of Dvorkin's model can also be clearly seen from the plots of Fig. 11 that at lower frequencies, the predicted values of $|E_{mix}|$ are lower than the expected values. Moreover, the predicted results of δ_{mix} do not show the expected decreasing trend. One of the possible reasons could be that the values of some input parameters, i.e. h_0 , n, a and v_b , should change with frequencies/temperatures rather than being constants over the whole frequency range. With this realization, in order to further understand the limitation of Dvorkin's model, the sensitivity analysis of the predicted stiffness on these parameters was conducted. In the following section, the results of the sensitivity analysis are presented.

6.2. Sensitivity analysis of predicted results to input parameters

In the previous analysis, the geometrical characteristics of the proposed PA mix's microstructure were considered to remain unchanged. However, this assumption may not be true because the external loads, i.e. the gravity and the applied load, may change the values of geometrical parameters (i.e. h_0 , n and a). With the increase of temperatures and the decrease of frequencies, the mortar layers become softer and easier to deform. Consequently, the distance between adjacent particles, i.e. the value of h_0 , is expected to decrease. Meanwhile, the change of h_0 also induces the change of n and a, which can be easily derived from Eqs. (B.9)–(B.12) and Eq. (B.16).

The above discussion indicates that the use of constant values of h_0 , n and a in the whole frequency range can be one reason for the poor performance of Dvorkin's model in Fig. 11. Additionally, a constant value of v_b was assumed in the predictions. As mentioned earlier, the value of v_b is frequency- and temperature-dependent. Therefore, it is also possible that the inaccuracy of the predictions results from the assumption of a constant value of v_b . With these

realizations, in this section, the values of h_0 and v_b were varied to check if the performance of the model can be improved. It is noted that sensitivity analysis was conducted by taking PA mix-A as an example.

6.2.1. The sensitivity of predicted E_{mix} on h_0

As discussed above, the value of h_0 is expected to decrease with the decrease of frequencies, therefore, three lower values of h_0 (<0.22 mm) were used to predict E_{mix} in the sensitivity analysis. Table 4 shows the calculated results of n and a for different values of h_0 using Eqs. (B.9)–(B.12) and Eq. (B.16), respectively. It can be seen that with the decrease of h_0 , the values of n and $\phi_{b_{-}c}$ decrease while the values of a and $\phi_{b_{-}b}$ increase. The reasons for the changes of $n, a, \phi_{b_{-}b}$ and $\phi_{b_{-}c}$ with the value of h_0 have been explained in the previous section (see Section 5.2).

Fig. 12 presents the predicted $|E_{\text{mix}}|$ -f and δ_{mix} -f curves using the values of h_0 , n and a in Table 4. It can be observed that with the decrease of h_0 , PA mixes become stiffer, which is reflected by an increase of predicted $|E_{\text{mix}}|$ and a decrease of predicted δ_{mix} . This can be explained by the fact that with the decrease of two particles' distance, the total volume of the binding mortar layers, which make the main contribution to the load-bearing capacity of the mix, increases. Meanwhile, when two particles become closer, their interactions become stronger. As can be derived from Eqs. (4) and (5), the stresses at the mortar-particle interface increase with the decrease of h_0 . When the interactions between adjacent particles become stronger, their stiffening effect on the mix becomes more significant.

Further investigations of the plots in Fig. 12 show that the shapes of the predicted $|E_{mix}|$ -f and δ_{mix} -f curves do not significantly change with different values of h_0 . At very low frequencies, even when h_0 is close to 0, the predicted values of $|E_{mix}|$ are still much lower than the experimental results. Moreover, the predicted values of δ_{mix} do not show a significant decrease. These observations indicate that the predictions are not likely to be accurate even if the values of the geometric parameters change.

Therefore, it can be concluded that the inaccurate assumption of geometric parameters is not the main reason to explain the poor performance of the model at high temperatures/low frequencies. With this realization, further investigation was conducted to evaluate the influence of another assumed parameter v_b on the predicted results.

6.2.2. The sensitivity of predicted E_{mix} on v_b

The predicted results of E_{mix} using different values of v_{b} are shown in Fig. 13. With the increase of v_{b} , the predicted values of $|E_{\text{mix}}|$ increase whereas the predicted values of δ_{mix} decrease. When the value of v_{b} reaches around 0.5, the predicted values of $|E_{\text{mix}}|$ become much higher than the experimental values whereas the values of δ_{mix} are far below. It can be postulated that if higher values of v_{b} are used at lower frequencies, the predicted values of | $E_{\text{mix}}|$ and δ_{mix} may match with the experimental results. In order to understand the effect of v_{b} on the predicted results of E_{mix} , and to figure out if increasing the value of v_{b} is a logical way to improve the accuracy of the predictions, the effect of v_{b} on the behavior of the binder layers was investigated.

In Dvorkin's model, the properties of the binder layers are described using Eqs. (4) and (5). The stiffness of the binder layer in the normal direction $S_{n,b}$ (2 $G_b(1-v_b)/(1-2v_b)$) is a function of both G_b and v_b . Fig. 14 shows the calculated results of $S_{n,b}/2G_b$ (or $(1-v_b)/(1-2v_b)$) when the value of v_b ranges from 0 to 0.5. It can be observed that when v_b is smaller than 0.45, the value of $S_{n,b}$ is at most 12 times of G_b . Since G_b is much lower than the stiffness of aggregates as well. In this case, the normal stiffness of the two bonded particles (F_n/δ_n) and the predicted stiffness of the mix are governed by the softer binder layer. Therefore, in Fig. 13, it was observed that when v_b is equal to 0.3 and 0.4, the predicted results of $|E_{mix}|$ and δ_{mix} follow the same shapes as $|G_b|$ and δ_b (see Fig. 10), respectively.

Table 4

Values of input parameters with the change of $h_{0.}$

Parameters	h_0/t	n	<i>a</i> (mm)	$\phi_{\mathbf{b}_{-}\mathbf{b}}$	ϕ_{b_c}
Values	0.58	11.5	1.67	9.4%	10.6%
	0.25	9.1	2.2	13.5%	6.5%
	0.125	8.8	2.35	14.9%	5.1%
	0	8.5	2.53	17.3%	2.7%



Fig. 12. The sensitivity of predicted $|E_{mix}|$ and δ_{mix} on $h_{0.}$



Fig. 13. The sensitivity of predicted E_{mix} on $v_{\text{b.}}$



Fig. 14. The sensitivity of $S_{n_b}/2G_b$ on v_b

On the contrary, when v_b is higher than 0.45, the value of S_{n_b} increases significantly. When v_b approaches 0.5, S_{n_b} becomes infinite, which indicates that the binder layer behaves as a rigid body. In this case, when F_n is applied, only aggregates deform and thus the predicted modulus of the mix relies on the properties of the aggregates. Since the aggregates are stiffer and behave as elastic materials, the predicted properties of the mix (see Fig. 13) tend to be stiff (high value of $|E_{mix}|$) and elastic (low values of δ_{mix}) as well.

From the above discussions, it can be concluded that the effect of v_b on predicted $|E_{mix}|$ and δ_{mix} is related to the assumption that with the increase of v_b , the mortar layer behaves like a rigid body. In this case, when two aggregate particles approach each other, the mortar layer does not have any deformation, while only aggregates deform. However, in a real PA mix, when v_b approaches 0.5 at high temperatures or low frequencies, it is expected that the mortar layer behaves more like a viscous material that easily deforms under loading, while the deformation of the stiff aggregates is much less significant. It is obvious that in these two cases, the properties of the mortar layer are different. Therefore, although the predictions with higher values of v_b may match with the experimental results numerically, the physical mechanism behind these predictions does not seem to be realistic.

6.3. Limitation of Dvorkin's model and possible explanations

Predicted results in Fig. 11 show that Dvorkin's model performs better in predicting the modulus of PA mixes at higher frequencies than CBMM, especially for mixes with densely packing aggregate particles; however, it still fails to provide accurate predictions at lower frequencies. Following sensitivity analyses further reveal that it is impossible (or unreasonable) to improve the accuracy of the predictions at lower frequencies by varying the values of input parameters.

The limitation of Dvorkin's model can be related to the assumption that in a bonded granular material, a load is always transferred through the mortar layers between adjacent particles. This assumption is clearly reflected by Eqs. (2) and (3), where the total deformation of a two-bonded particle system is the sum of the deformation of the particles and the deformation of the mortar layer. This assumption is suitable at low temperatures/high frequencies because, at these conditions, the mortar layer between two particles is stiff enough to transfer a high level of load from one particle to the other. Even if a direct particle-on-particle contact area forms at the same time, since the total particle-mortar contact area is expected to dominate over the particle-on-particle contact area, the overall stiffness of the mix is governed by the behavior of the mortar layers.

On the other hand, at high temperatures/low frequencies, the mortar layers are too soft to effectively transfer loads among adjacent particles. Comparing to the soft mortar layers, direct particleon-particle contacts are supposed to play a leading role in providing the load transfer capacity for the mix. In this case, the assumption in Dvorkin's model is not applicable anymore, and thus the predictions start to differ from the experimental results.

7. Conclusions

This study presented a methodology to use Dvorkin's model to predict the stiffness of PA mixes. The predicted results were compared to those from commonly used CBMM and the experimental values. In order to improve the accuracy of the predictions at lower frequencies, the sensitivity of the predicted results on geometrical characteristics of the mix and the Poisson's ratio of the binder layer was analysed. In the end, the limitation of Dvorkin's model was

highlighted. Based on the obtained results, the following conclusions can be drawn:

- In the proposed microstructure model, a PA mix was simulated as an assembly of identical spherical particles that are covered by a uniform thickness of mortar layers and bonded together by binding mortar layers.
- Geometric parameters were determined in a way that: the radius of the spherical particles was determined from the aggregate gradation; the thickness of the coating mortar layers was determined from the mortar content; the minimum distance between two adjacent particles and the coordination number were determined from air voids content, and the radius of the binding mortar layers was finally determined on the basis of the values of other parameters.
- Based on the proposed microstructure model, the stiffness of PA mixes was calculated in three steps. At first, the stiffness of the skeleton framework consisting of aggregate particles and the binding mortar layers was predicted using Dvorkin's model. Then the matrix's stiffness of an equivalent two-phase composite, whose stiffness was identical to that of the skeleton framework, was back-calculated using the Self-consistent model. At last, by adding the coating mortar layers in the matrix, the stiffness of PA mixes was calculated using the Self-consistent model again.
- Dvorkin's model describes a more realistic microstructure for PA type mixes. This model takes several geometrical characteristics of the aggregate particles and the mortar layers into account. Therefore, the utilization of Dvorkin's model is beneficial to better understand how the characteristics in the microscale affect the mechanical properties of a mix. Furthermore, in a wide range of high frequencies, Dvorkin's model can provide accurate predictions not only for loosely compacted PA mixes but also for PA mixes with densely packed aggregate particles.
- Dvorkin's models showed inaccurate predictions at lower frequencies: the predicted dynamic moduli were significantly lower than the experimental results, and moreover, the predicted phase angle did not show a decreasing trend. These poor performances cannot be significantly improved by varying the values of the geometric parameters. By varying the Poisson's ratio of the mortar layers, the predictions may match the experimental results numerically. However, the physical mechanism behind these predictions that the mortar layers behave like a rigid body does not seem to be realistic.
- The limitation of Dvorkin's model is related to the assumption that in a bonded granular material, a load is always transferred through the mortar layers between adjacent particles. This assumption is valid at higher frequencies, while at lower frequencies, the load is supposed to be mainly transferred through the direct particle-on-particle contacts.
- In future research work, the capability of Dvorkin's model to accurately predict the stiffness of PA mixes at higher frequencies will be further validated using PA mixes with different gradations and binder contents. Furthermore, Dvorkin's model will be modified to take into account the contribution from the direct particle-on-particle contacts. This modification may be achieved by using Hertzian contact theory to describe the mechanical responses of two contacting particles.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Data availability

The raw/processed data required to reproduce these findings cannot be shared at this time as the data also forms part of an ongoing study.

Appendix A. Numerical solutions to the displacement of binder layer

In this study, a numerical method was used to calculate V(x) and U(x). In the following paragraphs, the steps for solving V(x) in Eq. (9) are presented and the same method can be used to solve U(x) in Eq. (10). It is noted that the method used in this study was developed on the basis of the basic approaches for solving the Volterra integral equations of the second kind [40].

The calculation of V(x) can be made in three steps:

• Step1: choose the points where the value of V(x) is calculated. In the range from 0 to a, N + 1 discrete points of x with an identical distance between neighboring points, x_0, x_1, \ldots, x_N , are chosen. By substituting these points into Eq. (9), N + 1 equations are obtained, see Eq. (A.1). The following steps aim to solve these N + 1 equations to obtain the values of V(x) at different points of x.

$$\delta_{n} + V(x_{i}) = \Lambda_{n} \int_{0}^{\pi} d\varphi \int_{0}^{x_{i}\cos\varphi + \sqrt{a^{2} - x_{i}^{2}\sin^{2}\varphi}} \times \frac{V(\sqrt{x_{i}^{2} + s^{2} - 2x_{i}s\cos\varphi})}{h_{0} + \frac{x_{i}^{2} + s^{2} - 2x_{i}s\cos\varphi}{2R}} ds, \ i = 0, \ 1, \ ..., \ N$$
(A.1)

• Step 2: discretize the variables. The variable φ is identically discretized into N + 1 discrete points in the range from 0 to π , φ_0 , φ_1 , ..., φ_N . The discretization of *s* is dependent on the value of x_i . In order to solve the N + 1 unknowns of $V(x_i)$ in Eq. (A.1), it is required that no other unknowns are introduced. This means that for a certain value of x_i , the discretized values of *s*, s_0, s_1, \ldots, s_{ni} , should satisfy Eq. (A.2). Thus, the values of s_j can be calculated by using Eq. (A.3). Eq. (A.3) is a quadratic equation and the roots are given as Eq. (A.4). It is noted here that since the upper boundary of *s* is a function of x_i , the number of the discretized points of *s* (represented by n_i in Eqs. (A.3)–(A.4)) is dependent on the value of x_i .

$$V(\sqrt{x_i^2 + s_j^2 - 2x_i s_j \cos \varphi_k}) = V(x_j), \quad i = 0, \ 1, \ ..., \ N$$
(A.2)

$$x_i^2 + s_j^2 - 2x_i s_j \cos \varphi_k = x_j^2, \quad j = 0, \ 1, \ ..., \ n_i$$
(A.3)

$$s_j = x_i \cos \varphi_k \pm \sqrt{x_j^2 - x_i^2 \sin^2 \varphi_k}, \quad j = 0, \ 1, \ ..., \ n_i$$
 (A.4)

• Step 3: solve the integral equation. By substituting all the discretized points of φ and *s* into Eq. (A.1), the discretized form is written as Eq. (A.5). The values of $V(x_i)$ can be obtained by solving these N + 1 equations. In Eq. (A.5), ω_k^{φ} and w_j^s are the weights of the discrete points of φ_k and s_i , respectively. If the

trapezoidal rule is used for approximating the integral values, the values of ω_k^{φ} and w_j^s are equal to 1 for the middle points while they are equal to 0.5 for the points at the boundaries. h^{φ} is the distance between two neighboring points for φ , which is equal to π/N . h_{ij}^s is the distance between two neighboring points for *s*, which can be automatically known once the values of s_i are obtained.

$$\delta_n + V(x_i) = \Lambda_n \sum_{k=0}^N h^{\varphi} \omega_k^{\varphi} \sum_{j=0}^{n_i} h_{ij}^s \omega_j^s \frac{V(x_j)}{h_0 + \frac{x_i^2}{2R}}, \quad i = 0, \ 1, \ ..., \ N$$
(A.5)

Appendix B. Method to determine geometrical parameters

In the proposed microstructure in Fig. 5b, five geometric parameters, i.e. *R*, *t*, *h*₀, *n* and *a*, are required to predict the mix's effective modulus. In this appendix, the proposed methods to determine these parameters are presented. Once the geometric parameters are known, the volume fractions of the binding mortar layers $\phi_{b_{-}b}$ and the coating mortar layers $\phi_{b_{-}c}$ can be further determined. Thus, the proposed method to determine $\phi_{b_{-}b}$ and $\phi_{b_{-}c}$ is introduced as well.

B.1 Calculation of R

In Dvorkin's model, aggregate particles are modelled as identical spheres. However, in an asphalt mix, the size of the aggregate particles is generally not uniform but graded. Therefore, in order to specify the value of *R*, it is necessary to obtain a representative size of the aggregate particles in the mix. For this purpose, a commonly used mathematical method of averaging different sizes of graded aggregate particles was utilized [41], see Eqs. (B.1) and (B.2):

$$R = \sum_{i=1}^{N-1} \phi_i \cdot \frac{d_i + d_{i+1}}{2} \tag{B.1}$$

with

$$\phi_{i} = \frac{(P_{i+1} - P_{i})/\rho_{i}}{\sum_{i=1}^{N-1} (P_{j+1} - P_{j})/\rho_{i}}$$
(B.2)

where P_{i+1} and P_i are the percentages passing the sieve i + 1 and sieve i by the total weight of aggregates, respectively; ρ_i is the density of aggregates retained on sieve i; ϕ_i is the volume fraction of aggregate particles retained on sieve i; N is the aggregates' total grades by sieving, i.e., 0.063 mm, 2 mm, 5.6 mm, etc.; and d_{i+1} and d_i are the diameters of sieve i + 1 and sieve i, respectively.

B.2 Calculation of t

The value of *t* can be determined according to the binder content β in the mix, which is defined as

$$\beta = \frac{V_{\rm b}}{V_{\rm b} + V_{\rm p}} = \frac{\phi_{\rm b}}{\phi_{\rm p} + \phi_{\rm b}} \tag{B.3}$$

where $V_{\rm b}$ and $\phi_{\rm b}$ denote the volumes and the volume fraction of the mortar layers, respectively.

Assuming that individual particles are surrounded by mortar layers with identical thicknesses, the value of $V_{\rm b}$ can be calculated using Eq. (B.4).

$$V_{\rm b} = N_{\rm p} \cdot \frac{4}{3} \pi R^3 \left[\left(1 + \frac{t}{R} \right)^3 - 1 \right] \tag{B.4}$$

By substituting Eqs. (28) and (B.4) into Eq. (B.3), the value of β can be written as

$$\beta = 1 - \frac{1}{\left(1 + \frac{t}{R}\right)^3} \tag{B.5}$$

Since the value of β is known from the values of $\phi_{\rm b}$ and $\phi_{\rm p}$, the value of *t* can be computed from Eq. (B.5).

B.3 Calculation of h₀

As mentioned earlier, the value of h_0 is related to the compaction effort. Since the compaction effort can be reflected by the air voids content, the value of h_0 can be determined according to the air voids content of the mix.

Fig. B1 shows the relationship between the air voids content of the mix $\phi_{v,c}$ and the number of compaction cycles *N* during the compaction process. The value of $\phi_{v,c}$ was calculated according to the height of the specimens *H*, Eqs. (B.6)–(B.8).

$$\phi_{\rm v_c} = \frac{V_{\rm total} - V_{\rm mass}}{V_{\rm total}} \tag{B.6}$$

$$V_{\text{total}} = \frac{1}{4}\pi \times 150^2 \times H \tag{B.7}$$

$$V_{\rm mass} = \frac{W_{\rm mass}}{\rho_{\rm max}} \tag{B.8}$$

where V_{total} denotes the total volume of the specimen; V_{mass} and W_{mass} denote the volume and the weight of the real mass of the specimen; and ρ_{max} is the maximum density of the mix, which can be obtained from the density of each constituent.

Before any compaction effort is applied (when N = 0), the value of h_0 is expected to be equal to t. In this case, the corresponding value of $\phi_{v.c}$ is 0.22, see Fig. B1. With the increase of the compaction effort, the value of h_0 decreases and thus the value of $\phi_{v.c}$ decreases as well. After conducting a large amount of compaction effort, h_0 is expected to be gradually close to 0, and meanwhile, $\phi_{v.c}$ shows a gradually stable value of 0.13. It is highlighted here that in Fig. B1, there is no point where the tangent to the curve is horizontal; however, by fitting the curve using an exponential function ($\phi_{v.c} = ae^{-bN} + c$; where a, b and c are fitting parameters), it was obtained that the minimum value of $\phi_{v.c}$ was 0.13 (c = 0.13). Therefore, it was stated that $\phi_{v.c}$ gradually reached a stable value of 0.13.

Using the values of h_0 and $\phi_{v,c}$ in the above mentioned two extreme cases, the relationship between h_0/t and $\phi_{v,c}$ was assumed to be expressed as a simple linear function, see Fig. B2. From this



Fig. B1. Relationship between air voids content and number of compaction cycles.



Fig. B2. Relationship between h_0/t and air voids content.

relationship, once the air voids content of the PA mix specimen is known, the value of h_0/t can be obtained.

Two points are worth noting here:

- For PA mix specimens with different gradations and binder contents, the relationship between h_0/t and $\phi_{v,c}$ should be different since the packing conditions of the aggregates are different. Therefore, in order to build such a relationship for a certain type of PA mix, there is no need to compact more samples with different gradations.
- A linear function may not be the most suitable one to describe the relationship between h₀/t and φ_{v_c}. However, since h₀ is a geometric parameter in the microscale, it is difficult to determine other values of h₀ during the compaction process. Sophisticated image processing techniques may be a promising method to solve this issue; however, such work is beyond the scope of this study. Furthermore, the agreement between the experimental results and the predictions, which will be shown in the later section, verifies that using such a simple linear function is acceptable.

B.4 Calculation of n

For a PA mix system in Fig. 5b where spherical particles are covered by mortar layers, it is difficult to directly determine its average coordination number *n*. However, for a "dry" packing system consisting of only identical spheres, researchers [42] have proposed that its coordination number n_d can be estimated on the basis of the air voids content of the system ϕ_d , see Fig. B3. The rela-



Fig. B3. Relationship between the coordination number and the air voids content in a packing system of identical spheres.

tionship between n_d and ϕ_d can be developed by plotting their values in different regular packings, i.e. a simple cubic packing ($n_d = 6$, $\phi_d = 0.4764$), a tetragonal sphenoidal packing ($n_d = 8$, $\phi_d = 0.3954$), a pyramidal packing ($n_d = 10$, $\phi_d = 0.3019$) and a tetrahedral packing ($n_d = 12$, $\phi_d = 0.2595$). As shown in Fig. B3, the value of n_d shows an almost linear increase with the decrease of ϕ_d . Therefore, a regressed linear equation, see Eq. (B.9), can be given to describe the relationship between n_d and ϕ_d .

$$\phi_0 = -0.0374n + 0.6956 \tag{B.9}$$

On the basis of the relationship in Fig. B3, researchers [42] further developed a method to determine the average coordination number of a packing system where the binding mortar layers were included. Using a similar method, the value of n was determined in this study.

In order to determine the value of n, a reference packing system is introduced, see Fig. B4. In the reference system, individual particles have the same configurations as those in the PA mix in Fig. 5b. This indicates that the coordination number in the reference system remains the same as n. However, particles in the reference system are covered by mortar layers with a thickness of h_0 instead of a thickness of t. Due to the change of the mortar layers' thickness, the air voids content in the reference system changes to a value of ϕ_0 .

Since the minimum distance between adjacent particles in the PA mix system is 2 h_0 , particles covered with a mortar layer of h_0 in the reference system are supposed to just touch each other. Therefore, the arrangement of the mortar-coated particles in the reference system can be considered as the same as a dry packing system consisting of spheres with a uniform radius of $(R + h_0)$. Accordingly, the relationship between n and ϕ_0 can be described using the relationship in Eq. (B.9).

Total volumes of the particles and the mortar layers V_a in the reference system (see Fig. B4) can be calculated using Eq. (B.10), and thus the total volume of the reference system can be given as $V_a/(1-\phi_0)$. It is further assumed that more mortar materials are located around the particles in Fig. B4 to form the PA mix system in Fig. 5b. Since the total volume of the system cannot change, the relationship in Eq. (B.11) must satisfy.

$$V'_{\rm p} = N_{\rm p} \cdot \frac{4}{3} \pi (R + h_0)^3 \tag{B.10}$$

$$\frac{V_{\rm b} + V_{\rm p}}{1 - \phi_{\rm v}} = \frac{V_{\rm p}'}{1 - \phi_{\rm 0}} \tag{B.11}$$



Fig. B4. A reference packing system.

where $\phi_{\rm v}$ is the air voids content of the PA mix.

By substituting Eqs. (28), (B.4) and (B.10) into Eq. (B.11), the value of ϕ_v can be expressed as

$$\phi_{\rm v} = 1 - \frac{(1 - \phi_0) \cdot (1 + \tau)^3}{(1 + \gamma)^3} \tag{B.12}$$

where $\gamma = h_0/R$. By combining Eqs. (B.9) and (B.12), the value of *n* can be determined for a given value of ϕ_{γ} .

B.5 Calculation of a

It was mentioned in Section 3.1 that the binding mortar layer is composed of two parts. One part (with a radius of a_1) is formed because two mortar-coated particles overlap each other and the other part (with a radius of a_2) comes from the mortar that is squeezed out, see Fig. B5a. On the basis of this assumption, a mathematical relationship that the volume of the cylinder V_1 (bounded by the red line in Fig. B5b) is identical to that of the spherical cap V_2 (bounded by the green line) can be obtained:

$$H_1 \times \pi a^2 = \frac{\pi}{3} \times H_2^2 \times [3(R+t) - H_2]$$
(B.13)

where H_1 and H_2 are the heights of the cylinder and the spherical cap, respectively, and their values can be computed using Eq. (B.14) and Eq. (B.15), respectively.

$$H_1 = (R + h_0) - \sqrt{(R + t)^2 - a^2}$$
(B.14)

$$H_2 = (R+t) - \sqrt{(R+t)^2 - a^2}$$
(B.15)

By substituting Eqs. (B.14) and (B.15) into Eq. (B.13), the following relationship can be obtained:

$$\begin{split} \left[\sqrt{(1+\tau)^2 - \alpha^2} - (1+\gamma) \right] \alpha^2 &= \frac{1}{3} \left[(1+\tau) - \sqrt{(1+\tau)^2 - \alpha^2} \right]^2 \\ &\times \left[2(1+\tau) + \sqrt{(1+\tau)^2 - \alpha^2} \right] \end{split}$$

where $\alpha = a/R$. With the calculated values of *t* and h_0 from the previous steps, the value of *a* can be determined from Eq. (B.16).

B.6 Calculation of $\phi_{\mathbf{b}_{-}\mathbf{b}}$ and $\phi_{\mathbf{b}_{-}\mathbf{c}}$

As illustrated in Fig. B6, the volume of half of the binding mortar layer between two adjacent particles $v_{b_{-}b}$ can be calculated by subtracting the volume of a spherical cap from the volume of a cylinder, see Eq. (B.17). The total volume of the binding mortar lay-

ers in a PA mix $V_{b_{-b}}$ is equal to the value of $v_{b_{-b}}$ multiplying by the average coordination number *n* and the total number of the particles $N_{\rm p}$, see Eq. (B.18). Once the value of $V_{b_{-b}}$ is known, the value of $\phi_{b_{-b}}$ and $\phi_{b_{-c}}$ can be easily obtained on the basis of the total volume and the volume fraction of the mortar layers, see Eq. (B.19) and Eq. (B.20).

$$v_{b_b} = \pi \times a^2 \times (h_0 + H_3) - \frac{\pi}{3} \times H_3^2 \times (3R - H_3)$$
(B.16)

$$V_{b_b} = N_p \cdot n \cdot v_{b_b} \tag{B.17}$$

with

$$H_{3} = R - \sqrt{R^{2} - a^{2}}$$

$$\phi_{b_{b}} = \frac{V_{b_{b}}}{V_{b}} \cdot \phi_{b} \qquad (B.18)$$

$$\phi_{\mathbf{b}_{\mathbf{c}}} = 1 - \phi_{\mathbf{b}_{\mathbf{b}}} \tag{B.19}$$

where $V_{\rm b}$ is obtained from Eq. (B.4).

Appendix C. SC model for calculating matrix's stiffness of a twophase composite and the stiffness of a three-phase composite (as represented from Ref. [43])

The SC model assumes inclusions embedded into an infinite effective medium that possesses the same properties as the composite itself. Considering both the matrix and the inclusions are isotropic materials and the inclusions are spheres, the bulk modulus K and the shear modulus G of a N-phase composite can be expressed as

$$K_{\rm eff} = K_{\rm m} + \sum_{r=2}^{N} \frac{\phi_r (K_r - K_{\rm m}) (3K_{\rm eff} + 4G_{\rm eff})}{3K_r + 4G_{\rm eff}}$$
(C.1)

$$G_{\rm eff} = G_{\rm m} + \sum_{r=2}^{N} \frac{5\phi_r G_{\rm eff}(G_r - G_{\rm m})(3K_{\rm eff} + 4G_{\rm eff})}{3K_{\rm eff}(3G_{\rm eff} + 2G_r) + 4G_{\rm eff}(2G_{\rm eff} + 3G_r)}$$
(C.2)

where and the subscripts "m", "r" and "eff" denote the matrix phase, the inclusion phase r and the effective medium, respectively; ϕ denotes each phase's volume fraction.

For the calculation of the stiffness of the equivalent two-phase composite in Fig. 7a, Eqs. (C.1) and (C.2) can be reduced as Eqs. (C.3) and (C.4), respectively. It is noted that since the inclusion phase in the composite is air voids, the stiffness of the inclusion phase is zero.







$$K_{\rm eff} = K_{\rm m} + \frac{\phi_2(-K_{\rm m})(3K_{\rm eff} + 4G_{\rm eff})}{4G_{\rm eff}} \tag{C.3}$$

$$G_{eff} = G_m + \frac{5\phi_2 G_{eff}(-G_m)(3K_{eff} + 4G_{eff})}{9K_{eff}G_{eff} + 8G_{eff}^2}$$
(C.4)

where ϕ_2 is equal to the addition of ϕ_v and ϕ_{b_c} ; and K_{eff} and G_{eff} are identical to the moduli of the skeleton framework in Fig. 7a. Once the values of $K_{\rm eff}$ and $G_{\rm eff}$ are known, the values of $K_{\rm m}$ and $G_{\rm m}$ can be easily derived as:

$$K_{\rm m} = \frac{4K_{\rm eff}G_{\rm eff}}{4(1-\phi_2)G_{\rm eff} - 3\phi_2 K_{\rm eff}}$$
(C.5)

$$G_{\rm m} = \frac{G_{\rm eff}(9K_{\rm eff} + 8G_{\rm eff})}{3K_{\rm eff}(3 - 5\phi_2) + 4G_{\rm eff}(2 - 5\phi_2)} \tag{C.6}$$

Furthermore, in order to calculate the stiffness of PA mixes as a three-phase composite in Fig. 7b, Eqs. (C.1) and (C.2) can be rearranged as:

$$K_{\rm eff} = K_{\rm m} + \frac{\phi_2(K_2 - K_{\rm m})(3K_{\rm eff} + 4G_{\rm eff})}{3K_2 + 4G_{\rm eff}} + \frac{\phi_3(-K_{\rm m})(3K_{\rm eff} + 4G_{\rm eff})}{4G_{\rm eff}}$$
(C.7)

$$\begin{aligned} G_{\rm eff} &= G_{\rm m} + \frac{5\phi_2 G_{\rm eff}(G_2 - G_{\rm m})(3K_{\rm eff} + 4G_{\rm eff})}{3K_{\rm eff}(3G_{\rm eff} + 2G_2) + 4G_{\rm eff}(2G_{\rm eff} + 3G_2)} \\ &+ \frac{5\phi_3 G_{\rm eff}(-G_{\rm m})(3K_{\rm eff} + 4G_{\rm eff})}{9K_{\rm eff}G_{\rm eff} + 8G_{\rm eff}^2} \end{aligned} \tag{C.8}$$

where ϕ_2 and ϕ_3 are equal to $\phi_{\rm b}$ and $\phi_{\rm v}$, respectively; and K_2 and G_2 are equal to K_b and G_b , respectively. It is noted here that it is difficult to obtain explicit solutions for K_{eff} and G_{eff} from Eqs. (C.7) and (C.8); thus, a numerical method, i.e. the Newton-Raphson method, was applied in this study. The details about how to use this method can be found elsewhere [38].

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