Acceleration of the NVT-flash calculation for multicomponent mixtures using deep neural network models

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Abstract

Phase equilibrium calculation, also known as flash calculation, has been extensively applied in petroleum engineering, not only as a standalone application for separation process but also an integral component of compositional reservoir simulation. Previous research devoted numerous efforts to improve the accuracy of phase equilibrium calculations, which place more importance on safety than speed. However, the equation-of-state based flash calculation consumes an enormous amount of computational time in compositional simulation and thus becomes a bottleneck to the broad application of compositional simulators. Therefore, it is of vital importance to accelerate flash calculation without much compromise in accuracy and reliability, turning it into an active research topic in the last two decades.

With the rapid development of computational techniques, machine learning brings another wave of technology innovation. As a subfield of machine learning, the deep
neural network becomes a promising computational technique due to its great capacity to deal with complicated nonlinear functions, and thus attracts increasing attention from the academia and industry. In this study, we establish a deep neural network model to approximate the iterative flash calculation at given moles, volume and temperature, known as the NVT flash. A dynamic model designed for NVT flash problems is iteratively solved to generate data for training the neural network. In order to test the model’s capacity to handle complex fluid mixtures, three real reservoir fluids are investigated, including one Bakken oil and two Eagle Ford oils. Compared to previous studies that follow the conventional flash framework in which stability testing precedes phase splitting calculation, we incorporate stability test and phase split calculation together and accomplish both two steps by a single deep learning model. The trained model is able to identify the single vapor, single liquid and vapor-liquid state under the subcritical region of the hydrocarbon mixtures. A number of examples are presented to show the accuracy and efficiency of the proposed deep neural network. It is found that the trained model makes predictions at most 244 times faster than the iterative NVT flash calculation for the given cases and meanwhile preserves high accuracy.

**Introduction**

Phase equilibrium calculations are not only performed independently, for instance, in separation process, but also play a significant role in compositional simulation during which complex phase behaviors often take place and accompany with considerable mass transfer between hydrocarbon phases. As secondary recovery methods hardly bring further economical production from conventional reservoirs, enhanced oil recovery (EOR) techniques, like miscible flooding, are increasingly employed to improve oil productivity. The simulation of EOR processes requires accurate prediction of phase behaviors that could be modeled by the commonly-used equation of state (EOS), such as Peng-Robinson EOS\(^1\) or Soave-Redlich-Kwong EOS\(^2\), thus placing great importance on the EOS-based phase equilibrium
calculation. However, due to the large nonlinearity of phase equilibria problems, the popular iterative solution method for stability test and phase split calculation at given conditions (usually pressure, temperature and feed composition) is computationally expensive. It has been reported that up to 70% of computational time is consumed by flash calculation in compositional reservoir simulation. Although good initial approximations for phase splitting calculation can be obtained from stability test, the accuracy is preserved at the cost of efficiency, which is attributed to the repeated stability analysis, the high-dimension formulation and iterative algorithm itself. Since speed is one of critical concerns in current compositional simulator, speeding up flash calculation without too much compromise in accuracy and reliability is an active research topic in both academia and industry.

In reality, a hydrocarbon mixture may consist of tens to hundreds of species, which is far more than the number of components in numerical simulations. Thus, in order to accelerate phase equilibrium calculation, what first comes to mind is to lump the fluid mixture into a smaller number of pseudo-components without losing much accuracy of the EOS model.\(^3\) Clearly, the less the number of components is, the more computational time can be saved and meanwhile the more accuracy will be lost. Despite this, the speedup given by reducing the number of components is barely satisfactory, therefore stimulating a large batch of researchers to develop acceleration strategies for phase equilibrium calculation over the last two decades. Among them, the reduction method was first proposed by Michelsen\(^4\) in 1986, showing that only three equations need to be solved in two-phase flash problem under the Van der Waals mixing rule. The central idea of the reduced approach is to decrease the number of nonlinear equations and unknowns by taking advantage of the low-rank property of the binary interaction coefficient (BIC) matrix. However, his method, assuming all zero binary interaction coefficients, is severely circumscribed so that it’s hardly applicable to real reservoir fluids. Later, Jensen and Fredenslund\(^5\) extended the Michelsen’s research by introducing one non-hydrocarbon component for practical application. The nonzero BICs between the non-hydrocarbon and hydrocarbons yield five reduced variables, irrespective of the number
of components, but still limits the applicability of their approach. A generalized reduction theorem was then presented by Hendriks\textsuperscript{6}, which gets rid of the restriction on the BIC matrix. He reformulated phase equilibria problems with the transformed chemical potential, as a function of reduction variables. By introducing the truncated spectral method, Hendriks and Van Bergen\textsuperscript{7} simulated the two-phase equilibrium in the reduced space using Newton-Raphson iterations. Similarly, Pan and Firoozabadi developed reduction models for phase stability testing\textsuperscript{8} and phase equilibrium calculation\textsuperscript{9} based on the spectral theory of linear algebra. For stability analysis, they found the tangent-plane distance (TPD) function in the reduced space has smoother surface accompanied with a unique minimum, both of which significantly improve the robustness of their method. On the other hand, by extensively testing the conventional approach and reduced approach of the two-phase flash calculation, they concluded that the direct solution of the reduced model makes flash calculations more efficient and robust. In addition, Li and Johns\textsuperscript{10} developed their reduction model for rapid flash calculation by decomposing the BICs into two parts. The new phase split formulation includes at most six reduced parameters, either degenerating to the Michelsen’s reduction model with three reduced parameters when all BICs are zero or the five-reduced-parameter model of Jensen and Fredenslund when the nonzero BICs only exist between a single component and the other components. Moreover, their method avoids solving the nonlinear Rachford-Rice equation\textsuperscript{11}, a difficulty in phase splitting calculation, and meanwhile retains the accuracy of conventional approaches. Recently, Nichita and Gracia\textsuperscript{12} designed a reduced flash formulation with a new set of independent variables and error equations, leading to simpler expressions of the Jacobian matrix. Compared to the classical reduction methods, their approach exhibits better efficiency and meanwhile preserves the robustness. Instead of decomposing BIC matrices by the conventional spectral analysis approach, Gaganis and Varotsis\textsuperscript{13} introduced a new decomposition method to minimize the approximation errors of the energy parameter, which allows the phase equilibrium calculation to be performed at a given accuracy with fewer reduced variables. Even though all the above reduction meth-
ods have shown great potentials in acceleration of the flash calculation, recent research\textsuperscript{14–17} questioned its efficiency against the conventional method. No agreement has been reached at which points the reduced methods become faster than the conventional ones. However, the reduction model seems to be more applicable to mixtures with 6 to 15 components.

In addition, there are many other approaches to speed up flash calculation within the framework of compositional simulator. Wang and Stenby\textsuperscript{18} developed a non-iterative flash calculation algorithm on the assumption that thermodynamic equilibrium is instantaneously established at any time of the simulation. With flash calculations decoupled from the compositional flow, the iterative flash procedure can be linearly approximated so that the mole numbers of components in each phase are computed based on the solution of compositional flow between the previous and current time steps. There is no need to solve the Rachford-Rice equation for the phase saturation, thereby improving the efficiency of their method further. However, the reduction of CPU time varies case by case. Given that the repetitive phase stability test is time consuming, Rasmussen et al.\textsuperscript{19} proposed the shadow region method in order to bypass unnecessary stability testing based on the location of any given condition in the phase diagram. It is assumed that the pressure, temperature or compositions have small changes in a certain grid block between two consecutive time steps. In the two-phase region, stability test procedure is completely bypassed by initializing flash calculation with the equilibrium solution in the last time step. Furthermore, a one-sided stability test, initialized by the pre-stored non-trivial stability solution, replaces the full stability test within the shadow region, while outside the shadow region the distance to the critical point is used to determine whether stability analysis is skipped or not. Consequently, the computation time spent on flash calculation is significantly reduced. On the other hand, to accelerate the phase splitting calculation, Voskov and Tchelepi\textsuperscript{20–22} presented the Compositional Space Adaptive Tabulation (CSAT) method in which a tie-line table is constructed using the pre-stored phase split solutions to parameterize the compositional space. If the investigated composition is on the predetermined tieline or its extension, the phase equilibria solution
can be approximated on the basis of the tieline compositions; otherwise, the standard flash procedure is performed and an additional tieline will be added into the lookup table. It is noted that the accuracy and efficiency of the CSAT method heavily depends on the number of stored tielines, as well as the tolerance set to accept the feed composition. As the tieline table becomes larger, we gain more accuracy but lose more efficiency as well. Inspired by Voskov and Tchelepi’s research, Belkadi et al.\textsuperscript{23} suggested using the Tieline Distance-Based Approximation (TDBA) method to approximate flash solutions in the two-phase region. In their approach, the shortest distance from the new feed composition to the tieline in the same gridblock is used to determine if the rigorous flash solution is approximated, which minimizes the access and build-up process of the tieline table. Recently, Yan et al.\textsuperscript{24} accelerated phase behavior calculations by interpolating initial approximations from the compositional space constructed, prior to the flash calculation, for reservoir fluids with both fixed and dramatically varying composition. For those hydrocarbon mixtures with significant compositional change during simulation, an additional pseudo key component is introduced, together with pressure, to control the phase behavior prediction. Although their approach exhibits an enhanced performance under the Rachford-Rice preconditioning, the resolution of the compositional spaces constructed offline play a critical role in such kind of methods as expected. In addition, Wu et al.\textsuperscript{25} utilized the sparse grid surrogate model to approximate flash results during the two-phase compositional simulation. Under a similar interpolation philosophy, a remarkable speedup of compositional simulation was achieved, but several intrinsic defects restrict the applicability of their method. Subsequently, Wu and Chen\textsuperscript{26} extended their previous work by approximating the equilibrium solution directly from the sparse grid instead of the surrogate model. The introduction of the "layer" concept and array structure makes the sparse grid construction efficient and inexpensive. However, two issues, including the violation of saturation boundedness and inaccurate detection of the single-phase region, need to be resolved for reliable application of sparse grid methods in phase equilibrium calculation.

Apart from the aforementioned accelerating strategies, other alternatives, like negative
flash calculation\textsuperscript{27,28} or linear transformation of the Rachford-Rice equation\textsuperscript{29}, can be used to improve the efficiency of flash calculation as well. All the above speedup approaches are only the tip of iceberg and few of them can avoid iteratively solving phase equilibria problems. In addition to the iterative nature of flash algorithms, the increasing demand to high-resolution reservoir models, usually with millions of grid blocks, also sharply increases the computational costs of phase equilibrium calculations during compositional simulation. Thus, it is imperative to perform both phase stability test and phase split calculation in a fast, non-iterative and robust way. With the rapid development of computing techniques, machine learning has become a powerful tool to deal with those sophisticated and time-consuming problems. Since 2012, due to the breakthrough of AlexNet\textsuperscript{30}, deep learning, a subset of machine learning, made a revolutionary progress in image recognition, language processing, artificial intelligence, and so forth. The great potential of machine learning has drawn close attention from the petroleum industry, and remarkable efforts have been made in many aspects, such as reservoir characterization\textsuperscript{31}, development planning optimization\textsuperscript{32} and permeability estimation.\textsuperscript{33} Furthermore, a number of studies have been devoted to speed up both phase stability test and phase split calculation using the machine learning approach. Gaganis and Varotsis\textsuperscript{34} designed a stability discriminating function, which yields the same sign and zero points as the conventional tangent plane distance (TPD) approach, to eliminate the time-consuming iterations in stability analysis. Under the framework of Support Vector Machine (SVM)\textsuperscript{35–37}, phase stability test is treated as a binary classification problem and has an explicit expression on the basis of pressure, temperature and feed composition. For phase split calculation, they developed regression models using the artificial neural network (ANN) to predict the conventional equilibrium coefficients\textsuperscript{38} and later the prevailing reduced variables\textsuperscript{39}, in order to approximate the closed-form solution of the original flash formulation and the reduced one respectively. Recently, Gaganis\textsuperscript{40} presented a new classification technique using two simplified discriminating functions to achieve rapid stability determination without much compromise in accuracy to depict the phase boundary. Instead of using
SVM, Kashinath et al.\textsuperscript{41} utilized two relevance vector machine (RVM) classifiers for stability analysis. The supercritical classifier identifies the supercritical state at the given condition, while the subcritical classifier predicts the number of stable phases. Similar to Gaganis and Varotsis’s approach, they estimated the equilibrium coefficients for the phase split problem using the regression model established on the basis of ANNs. Unlike the aforementioned works, Wang et al.\textsuperscript{42} built machine learning models based on ANNs for both stability test and phase split calculation. The ANN model for stability testing predicts the saturation pressure, which is used for stability determination in comparison with the system pressure. The other ANN model provides initial mole fractions and equilibrium coefficients for phase split calculations. In addition, Li et al.\textsuperscript{43} approximately solved the vapor-liquid equilibria problem using a deep learning model. Different from the aforementioned works, they directly estimated the equilibrium compositions rather than the equilibrium coefficients that work as the intermediate variables. A variety of effects are investigated, such as the number of hidden layers, the number of nodes per layer and the type of activation function, on the performance of the deep neural network, which helps them optimize their model to improve efficiency and accuracy. However, as a shortcoming of their work, they only discussed the cases where both vapor and liquid phases are present.

In this study, instead of using the conventional NPT-formulation, the phase equilibrium problem is formulated at constant moles, volume and temperature (the so-called NVT flash). A dynamic model is designed for this isothermal-isochoric phase equilibria problem and iteratively solved by Newton-Raphson methods with line search. To validate the capacity of our dynamic model, as well as the deep learning model, for handling complex fluid mixtures, three realistic reservoir fluids are investigated: one from Bakken shale formation includes 5 components and the other two from Eagle Ford shale formation consist of 8 and 14 components respectively. With the training data provided by the iterative flash calculation, we develop a deep neural network to predict phase compositions at different overall concentration and temperature with fixed feed composition, enabling it to deal with depletion processes. To
overcome the overfitting issue inherent in neural networks, the dropout technique is applied in combination of a regularized loss function. In addition, batch normalization significantly reduces the time spent on training the network model, thus enabling us to construct a more complex multi-level structure for better approximation. The network configuration is designed based on our previous research. Different from the preceding studies, one key effort of this study is to incorporate stability test and phase split calculation together so that both steps in the conventional flash framework can be accomplished by a single deep neural network. An independent variable is introduced in the training process to determine the number of stable phases. Meanwhile, the trained model identifies the fluid state (e.g., single vapor, single liquid or vapor-liquid) under the subcritical region. We present a number of numerical examples to validate the accuracy and efficiency of the proposed deep learning model.

The remainder of this paper is organized as follows. In the following section, we formulate the dynamic model for the NVT flash problem in the bulk phase and compare our simulation results with the published ones. In Section 2, the deep learning algorithm is elaborated in detail with some techniques to enhance its performance. We discuss the prediction accuracy of the proposed deep neural network and compare its efficiency with the iterative flash calculation in Section 3. At the end, we make our conclusions in Section 4.

1. NVT-flash calculations

Compared to the classical NPT-flash calculation, the NVT flash exhibits some advantages and attracts a lot of attention from researchers in recent years. Without inverting the equation of state, the NVT-formulation has a unique solution and thus eliminate the root-selection procedure that usually takes place in NPT flash problems. Additionally, volume of pure substances under saturation pressure cannot be uniquely determined using the NPT formulation, since all states (two phases, single vapor or single liquid phase) of a pure sub-
stance share the same pressure on the phase boundary. Similar behavior has been observed in multicomponent mixtures with three or four phases as well. Despite the fact that the NPT flash is the most commonly used flash technique in compositional simulators, numerous efforts have been made to improve the performance of the NVT flash calculation and also extend its applications. Jindrová and Mikyška developed a fast and robust NVT-flash algorithm for two-phase equilibria problems by directly minimizing the total Helmholtz free energy. Subsequently, they extended their research to multiphase systems and investigated the equilibrium problem involving aqueous phase through the Cubic-Plus-Association EOS. Similarly, Nichita formulated the NVT flash problem as unconstrained minimization of the Helmholtz free energy as well. Unlike Jindrová and Mikyška’s method, volume is treated as a dependent variable on mole numbers, leading to a "PT-like" iteration framework for the NVT flash calculation. To model the dynamic process from any non-equilibrium state to the equilibrium state, Kou et al. established evolution equations for mole numbers and volume, which was solved by a well-designed energy-stable numerical algorithm. In addition to the aforementioned bulk phase flash problems, the confined phase behaviors at constant moles, volume, temperature have been studied by taking into account adsorption, capillary pressure, or confinement effect resulting from the interaction between fluid molecules and pore walls. In this study, we restrict our discussion to the bulk phase equilibrium of two-phase systems in which the fluid mixture is described by the Peng-Robinson EOS.

1.1. The dynamic model for bulk phase equilibria

Let us assume a fluid mixture of components stays in the vapor-liquid equilibrium at the specified moles, volume and temperature. Thus, the total Helmholtz free energy of this system can be expressed as

\[ F = f(n^G) V^G + f(n^L) V^L \]
where \( \mathbf{n}^\alpha = [n_1^\alpha, \ldots, n_M^\alpha]^T \) and \( V^\alpha \) denote the molar density and volume of phase \( \alpha (\alpha = G, L) \), respectively. The Helmholtz free energy density \( f (\mathbf{n}) \) has the following form

\[
f (\mathbf{n}) = RT \sum_{i=1}^M n_i (\ln n_i - 1) - nRT \ln (1 - bn) + \frac{a(T)n}{2\sqrt{2b}} \ln \left( \frac{1 + (1 - \sqrt{2}) bn}{1 + (1 + \sqrt{2}) bn} \right)
\]  

(2)

In eq 2, \( n = \sum_j n_j \) is the overall molar density, \( R \) is the ideal gas constant and \( T \) is the temperature. The Peng-Robinson parameters \( a \) and \( b \) are functions of the molar composition. Considering the total mole numbers \( \mathbf{N}^t = [N_1^t, \ldots, N_M^t]^T \) and total volume \( V^t \) are fixed, the following mole and volume constraints have to be satisfied whenever both vapor and liquid phases are present in the system

\[
N_i^G + N_i^L = N_i^t \quad i = 1, \ldots, M
\]  

(3)

\[
V^G + V^L = V^t
\]  

(4)

where \( N_i^\alpha \) represents the mole number of component \( i \) in phase \( \alpha \).

In any two-phase system, we can arbitrarily select the mole composition and volume of one phase as primary variables and compute the counterparts of the other phase by the aforementioned mole and volume constraints. Here the mole composition and volume of the vapor phase, denoted by \( \mathbf{N}^G \) and \( V^G \), are chosen as the primary variables so that the Helmholtz free energy is reduced as \( F (\mathbf{N}^G, V^G) \). The partial derivatives of the Helmholtz free energy in eq 1 with respect to \( N_i^G \) and \( V^G \) yield

\[
\frac{\partial F (\mathbf{N}^G, V^G)}{\partial N_i^G} = \mu_i (\mathbf{n}^G) - \mu_i (\mathbf{n}^L) = \mu_i^G - \mu_i^L
\]  

(5)

\[
\frac{\partial F (\mathbf{N}^G, V^G)}{\partial V^G} = p (\mathbf{n}^L) - p (\mathbf{n}^G) = p^L - p^G
\]  

(6)

where \( \mu_i^\alpha \) and \( p^\alpha \) are the chemical potential of component \( i \) and pressure in phase \( \alpha \), respectively. By applying the chain rule, the time derivative of the total Helmholtz free energy
\( F(N^G, V^G) \) can be derived as follows

\[
\frac{\partial F}{\partial t} = \sum_{i=1}^{M} \frac{\partial F}{\partial N^G_i} \frac{\partial N^G_i}{\partial t} + \frac{\partial F}{\partial V^G} \frac{\partial V^G}{\partial t} \\
= \sum_{i=1}^{M} (\mu_i^G - \mu_i^L) \frac{\partial N^G_i}{\partial t} + (p^L - p^G) \frac{\partial V^G}{\partial t}
\] (7)

The mole and volume evolution equations are formulated based on Onsager’s reciprocal principle, shown as below, to characterize the dynamic bulk phase equilibrium process

\[
\frac{\partial N^G_i}{\partial t} = \sum_{j=1}^{M} \psi_{i,j} (\mu_j^G - \mu_j^L) + \psi_{i,M+1} (p^L - p^G)
\] (8)
\[
\frac{\partial V^G}{\partial t} = \sum_{j=1}^{M} \psi_{M+1,j} (\mu_j^G - \mu_j^L) + \psi_{M+1,M+1} (p^L - p^G)
\] (9)

According to the second law of thermodynamics, the total Helmholtz free energy in a closed system is supposed to dissipate with time, implying the Onsager coefficient matrix \( \Psi = (\psi_{i,j})_{i,j=1}^{M+1} \) shall be negative definite. Following the similar strategy in Ref. 50, we compute the Onsager coefficient matrix based on the equilibrium conditions. A revised modified Cholesky factorization is applied to ensure its negative definiteness, see Ref. 53 for more details. It is worth mentioning that this well-designed coefficient matrix helps to achieve the persistent dissipation of the Helmholtz free energy, but also simultaneously solve all evolution equations as a whole system in an iteration approach. When the Helmholtz free energy is minimized, the investigated system reaches the equilibrium state, including the chemical equilibrium (\( \mu_i^G = \mu_i^L \)) and the pressure equilibrium (\( p^G = p^L \)).

1.2. Numerical experiments

To perform the NVT flash calculation, each phase split calculation is initialized by the stability test in this study. Detailed implementation procedures of the bulk phase stability testing can refer to either the work of Mikyška and Firoozabadi\textsuperscript{54} or the work of Kou
and Sun\textsuperscript{49} with negligible capillary pressure. To construct an energy-stable time-marching scheme, which preserves the dissipation of the Helmholtz free energy, the convex splitting technique is employed to decompose the Helmholtz free energy density and chemical potential. In addition, we use a semi-implicit scheme\textsuperscript{55} to discretize the mole and volume evolution equations. Specifically, the convex parts of the chemical potential and Helmholtz free energy density are treated implicitly while the concave parts are treated explicitly. The resultant nonlinear system is solved by the Newton-Raphson method with line search. We terminate the Newton iteration if either the maximal number of iterations (500) or the Euclidean norm of the directional vector less than $10^{-8}$ is satisfied.

In order to validate the accuracy of the proposed dynamic model, three reservoir fluid samples are tested: the Bakken oil consists of 5 components and two Eagle Ford oils have 8 and 14 components. For brevity, the Eagle Ford oils are named by EagleFord1 for the eight-component sample and EagleFord2 for the other one with fourteen components. The molar composition and compositional properties for the Bakken, EagleFord1 and EagleFord2 oils are included in the Supporting information (see Table S1–S3). Binary interaction coefficients for these three samples can be found in Ref. 56–58 respectively. To achieve a smooth phase envelope, we uniformly discretize the computational domain of the overall molar density and temperature into a $201 \times 201$ grid. For the Bakken oil, the investigated molar density varies from 10 to 10000 mol/m$^3$ and the temperature ranges from 300 to 850 K. Moreover, the overall concentrations of EagleFord1 and EagleFord2 samples are $[10, 12000]$ and $[10, 10000]$ mol/m$^3$, while both of them have the same temperature interval $T \in [260, 850]$ K. Figure 1 shows the computed phase envelopes for the Bakken oil (top left), the EagleFord1 oil (top right) and the EagleFord2 oil (bottom). It can be seen that the equilibrium solutions of the dynamic model agree with the published results in Ref. 56–58 very well. In the following, we use these equilibrium results of NVT flash calculation, especially the mole fractions of vapor and liquid phases, to train our deep learning model.
Figure 1: Bulk phase envelopes for: the Bakken oil (top left), the EagleFord1 oil (top right) and the EagleFord2 oil (bottom).

2. Deep learning algorithm

Artificial neural networks are computational models that are able to unearth the underlying correlations and features of input data and process them in a manner analogous to the biological nervous system. Such a characteristic enables the ANN model great problem-solving capability. Various applications have been developed on the basis of ANNs, which are
performed in either supervised or unsupervised approach, such as machine learning or pattern recognition. Unlike the unsupervised ANN model, the supervised machine learning tries to seek a function that can be used to predict the values of desired variables from input data under a given accuracy. In this study, instead of using shallow neural networks, a deep neural network, which consists of multiple nonlinear transformation layers, is applied to replace the iterative NVT-flash algorithm in order to accelerate phase equilibrium calculations. Figure 2 displays the schematic diagram of the proposed deep neural network, which processes the input data, including compositional properties (e.g. critical pressure $P_c$, critical temperature $T_c$, acentric factor $\omega$), overall mole fractions $z = [z_1, \ldots, z_M]^T$, overall molar concentration $C$ and temperature $T$ in a sequential approach. Each activation layer transforms the output of the preceding layer and yields results as input to the following one. This hierarchical structure enables the deep neural network to extract useful features and patterns layer by layer and then learn complicated correlations behind the enormous data volume. Benefiting from the powerful capacity of the deep learning model, we accomplish both stability test and phase split calculation by a single network model so that the number of phases is determined free of additional stability test. This differs from the recent research on the acceleration of flash calculation by machine learning models, in which the stability test is preserved to determine if the investigated fluid is stable or not as the prevailing flash framework does. As a result, the output of the regression layer in the proposed neural network model includes three parts: the number of phases ($N$), the mole fraction of vapor components $X_i$ and the mole fraction of liquid components $Y_i$.

The performance of a neural network model can be evaluated based on the loss, which is defined as the difference between the model output and the ground truth. Here we use the mean square error as the loss function, evaluated in each iteration, to describe the deviations of the model prediction from the solution of iterative flash calculation. However, overfitting, as the inherent issue of neural networks, gravely affects the performance of the deep learning algorithm, which manifests that the trained model exhibits excellent performance based on
the training data while yields terrible predictions for its further application. This inconsistent behavior is attributed to the numerous parameters of the model waiting for optimization and insufficient training data, thus making the model over-parameterized. In order to protect the deep neural network model from the overfitting issue, additional constraints are imposed on the parameters to reduce the freedom of the model. Generally, if the model is overfitted, the norm of the weight parameters will become very large. To avoid the overfitting problem, an additional constraint is added on the norm of weight parameters to penalize the large weights. In practice, we introduce a regularization term, related to the norm of the weights, in the loss function, which yields

\[ L = \frac{1}{N} \sum_{n=1}^{N} \| o - \hat{o} \|^2 + \lambda \| W \|_2^2 \]  

(10)

where \( N \) is the number of the training data, \( o \) is the model output, \( \hat{o} \) is the observation value, \( W \) is the weight parameters, and \( \lambda \) is the regularization coefficient for weight penalization.

In addition to the regularized loss function, we also use the dropout technique to resolve
the overfitting problem, which reduces the freedom of the deep neural network model by abandoning certain nodes and connections during the training stage. In particular, with a specified threshold probability \( p (0 < p < 1) \), each node of a layer is evaluated independently and retained if its probability is less than the threshold value; otherwise, the node will be removed. The connections belonging to those disregarded nodes are removed as well, therefore yielding a reduced neural network for training. After this, the full model is recovered by inserting the removed nodes and connections back and enters the next training cycle.

In order to design a robust and efficient neural network, it is crucial to properly initialize the weight parameters since they have significant influence on the convergence rate and performance of the deep learning algorithm. If the initial weights are underestimated, the variance of the input data will drop to a very small value through some intermediate transformations, which reduces the model complexity and damage its performance. On the other hand, the overestimated weight parameters increase the variance of the input rapidly, making the gradient either vanishing or exploding and eventually failing to train the model.

In this study, the weight parameters are initialized following the Gaussian distribution. To control the variance during the training process, we initialize the weights with a variance \( \delta \) and make the output variance always same to the input variance on the same activation layer. For example, in Figure 3, the variance of \( y \) is expected to expand as follows

\[
\operatorname{var}(y) = \operatorname{var}(w_1 \times a_1 + w_2 \times a_2 + \ldots + w_n \times a_n + b)
\]

\[
= \operatorname{var}(w_1) \times \operatorname{var}(a_1) + \operatorname{var}(w_2) \times \operatorname{var}(a_2) + \ldots + \operatorname{var}(w_n) \times \operatorname{var}(a_n)
\]

\[
= n \times \operatorname{var}(w_i) \times \operatorname{var}(a_i)
\]

(11)

with the identical distribution assumption for all \( w_i \) and \( a_i \). Since we want \( \operatorname{var}(y) \) equal to \( \operatorname{var}(a_i) \), the following equality has to be satisfied

\[
 n \times \operatorname{var}(w_i) = 1.
\]

(12)
Therefore, the weights of each layer are initialized following the Gaussian distribution with the variance of $1/n$, where $n$ is the number of weights in that layer. This initializer is known as the Xavier initializer.

It has to be mentioned that training a deep neural network is time-consuming due to massive parameters waiting for optimization over all the hidden layers. In addition, some undesirable properties of the multi-level model, as well as the input data of different scale at each activation layer, slow down the training process. Among others, the varying distribution of the activations play a critical role in the slow convergence of training deep neural networks, since each layer has to adjust itself to a new distribution in each training step. To speed up the learning process, batch normalization is employed to adjust and scale the outcomes of the preceding layer, which can be regarded as a preprocessing step for each activation layer. Consequently, a significant improvement of the training speed is achieved, which enables to train more complicated models in a reasonable amount of time.

In this study, we employ a deep neural network model to approximate the iterative NVT-flash calculation using TFlearn on a MacBook Pro laptop with Intel Core i7 processor. A simplified flowchart of the working process in each node of the proposed model is presented in Figure 3. Let $a_i$ and $y_i$ denote the input and output of the $i$th activation layer, respectively. The output of the $i$th layer is given by

$$y_i = f_i(W_i \times a_i + b_i), \quad (13)$$

where $W_i$ is the weight, $b_i$ is the bias, and $f_i$ is the activation function of the $i$th layer. The neural network in Figure 2, for instance, with three activation layers, can be expressed as

$$o = f_3(W_3 \times f_2(W_2 \times f_1(W_1 \times x_1 + b_1) + b_2) + b_3). \quad (14)$$

As shown in eq 14, the activation function is the source of the nonlinearity in neural networks, which helps the deep learning algorithm approximate complex functions and gives it great
problem-solving capability. There are a variety of activation functions, including Rectified linear unit (ReLU), Parameteric rectified linear unit (PReLU), TanH, Sigmoid, Softplus, Softsign, Leaky rectified linear unit (Leaky ReLU), Exponential linear unit (ELU) and Scaled exponential linear unit (SELU). Among them, the most commonly used activation functions are ReLU

\[
f(x) = \begin{cases} 
0, & \text{if } x < 0 \\
 x, & \text{if } x \geq 0 
\end{cases}
\]

\hspace{1cm} (15)

and Sigmoid

\[
\sigma(x) = \frac{1}{1 + \exp(-x)} .
\]

\hspace{1cm} (16)

3. Results and Discussion

The training data for the deep learning model are provided by the NVT flash calculation. Three real reservoir fluids are investigated, including the five-component Bakken oil,
eight-component EagleFord1 oil and fourteen-component EagleFord2 oil. The compositional parameters for each reservoir fluid are presented in the Supporting Information. Following the theory and practice described in Section 2, a deep neural network is established with 5 activation layers, each of which contains 100 nodes, and totally 4000 iterations. "ReLU" is chosen as the activation function. It is worth mentioning that the performance of this network configuration has been validated in Ref. 43. To investigate the effect of data size on the performance of the deep neural network model, we calculate equilibrium results of the NVT flash for the EagleFord1 oil on the same computational domain with $51 \times 51$, $71 \times 71$, $101 \times 101$, $151 \times 151$, $201 \times 201$ and $301 \times 301$ uniform grids. In addition, the results of the Bakken oil and EagleFord2 oil are computed on the specified concentration and temperature intervals, which are uniformly divided into $301 \times 301$ grids. All the eight data sets are used to train the proposed neural network, and the efficiency and accuracy of the trained model are tested. One key effort of this study is to investigate the possibility in achievement of both stability test and phase split calculation by a single neural network model, which is different from the conventional two-step framework that all the preceding research follows based on machine learning models.

3.1. Deep Learning Model Training

As shown in Figure 2, the compositional properties of fluid components, overall molar concentration and temperature are used as the input, and the proposed deep neural network predicts mole fractions of components in both vapor and liquid phases. The key parameters of the model are the weights of each activation layer, which control the model prediction under the given input data. At the beginning, those weights are initialized randomly, implying that the model initially yields useless results. To approximate the NVT flash calculation by the deep learning model, we optimize the weight parameters to fit the equilibrium mole fraction of vapor and liquid components. In the following, 90% of the data are used to train our network model, while the remaining 10% data are used for validation unless otherwise
Table 1: Performance of the deep learning model for different data samples.

| source          | $N_{\text{train}}$ | $N_{\text{test}}$ | $t_{\text{train}}$ (s) | $t_{\text{test}}$ (s) | $\varepsilon_a$ | $\varepsilon_r$ |
|-----------------|---------------------|---------------------|-------------------------|-------------------------|-----------------|-----------------|
| Bakken (301 × 301) | 81540               | 9061               | 2352                    | 9.2                     | 0.02362         | 0.02885         |
| EagleFord1 (51 × 51) | 2340               | 261               | 42                      | 4.1                     | 0.02946         | 0.03582         |
| EagleFord1 (71 × 71) | 4536               | 505               | 134                     | 5.9                     | 0.01930         | 0.02410         |
| EagleFord1 (101 × 101) | 9180              | 1021              | 235                     | 6.3                     | 0.01378         | 0.01723         |
| EagleFord1 (151 × 151) | 20520             | 2281              | 512                     | 7.1                     | 0.01301         | 0.01647         |
| EagleFord1 (201 × 201) | 36361             | 4040              | 1232                    | 7.9                     | 0.01261         | 0.01580         |
| EagleFord1 (301 × 301) | 81540             | 9061              | 2468                    | 8.9                     | 0.01684         | 0.01964         |

Table 1 presents the training data size ($N_{\text{train}}$), testing data size ($N_{\text{test}}$), training time ($t_{\text{train}}$) and testing time ($t_{\text{test}}$) of the deep neural network with different data sets. Here $t_{\text{test}}$ represents the time that the trained model spent on estimating equilibrium mole fractions for the flash problem of same data size. For instance, it takes 7.9 seconds for the trained model to predict the mole fractions of the EagleFord1 oil on a 201 × 201 grid. In addition, the mean absolute error ($\varepsilon_a$) and relative error ($\varepsilon_r$) are presented in Table 1 as well. Clearly, for the EagleFord1 oil, as the number of input data becomes larger, the training time significantly increases, while the testing time doesn’t change too much. Furthermore, we observe that both absolute and relative prediction errors continue to decrease with the data size increasing. Under the same data volume, it seems the more components are involved, the larger prediction error the deep neural network model exhibits. However, the Bakken oil makes an exception and yields greater error than the Eagle Ford oils. This might be attributed to the underneath correlations between different components differ from the investigated fluid mixtures so that the trained network model yields different accuracy. Essentially, the composition of the Bakken oil is quite different from the compositions of two Eagle Ford samples, the latter of which exhibit some similarities to some extent. This may explain our observation disagrees with the expectation. Table 2 compares the computational time of NVT flash calculations to the testing time of the deep neural network for the EagleFord1 oil.
with different data size. It can be seen that the testing time is much less than the computational time of flash calculations. When the data size reaches to $301 \times 301$, the trained model makes predictions 244 times faster than the iterative flash calculation.

Table 2: Comparison of the computational time of the iterative flash calculation to the testing time of the deep neural network for the EagleFord1 oil with different data size.

| source                  | $t_{\text{flash}}$ (s) | $t_{\text{test}}$ (s) |
|-------------------------|-------------------------|-------------------------|
| EagleFord1($51 \times 51$) | 61                      | 4.1                     |
| EagleFord1($71 \times 71$) | 122                     | 5.9                     |
| EagleFord1($101 \times 101$) | 237                     | 6.3                     |
| EagleFord1($151 \times 151$) | 547                     | 7.1                     |
| EagleFord1($201 \times 201$) | 1021                    | 7.9                     |
| EagleFord1($301 \times 301$) | 2100                    | 8.6                     |

Figure 4 shows the loss function decreases with iterations, indicating the predictions of the trained model are approaching to the results of the NVT flash calculation. The inset figure displays the local variation of the total loss function after a sharp decrease within the first 100 iterations, which is indistinguishable in the main figure. It is observed that the difference between deep learning predictions and iterative flash solutions continues to decrease, but the decline rate significantly slows down after about 40 or 50 iterations. For the EagleFord1 oil, the increasing data size, from $51 \times 51$ to $201 \times 201$, helps to accelerate the reduction of the loss function. However, such a speedup effect vanishes with more data used to train the model. In comparison, the loss function of the EagleFord2 oil exhibits a faster decline rate with the same data size (e.g. $301 \times 301$) to the EagleFord1 oil until it reaches a certain threshold, below which the loss function remains almost steady.
3.2. Phase Number Characterization

Previous research follows the conventional framework of flash calculation, which consists of two separate stages: stability test and phase split calculation. In contrast, one key effort of this study is training the deep neural network to automatically determine the number of phases without additional stability testing. Under this guideline, the proposed neural network model completes both stability test and phase split calculation simultaneously, and predicts the number of phases together with the molar compositions of vapor and liquid phases, as shown in Figure 2. To validate the prediction accuracy of our model on phase stability problems, we compare the number of phases computed by the deep learning
model and iterative flash calculation. Figure 5 displays the phase number of the EagleFord1 oil as a function of the temperature at two overall concentrations 1988 mol/m³ (top) and 2648 mol/m³ (bottom), respectively. Note that we only plot the results slightly across the phase boundary in Figure 5. Overall, our deep learning model successfully captures the phase transition process from the vapor-liquid region to the single vapor region. The single-phase state, either single vapor or liquid, is determined based on the fact that the mole fractions of components in the existing phase should approach to overall mole fractions. Specifically, if the mixture is a single vapor (liquid) phase at the given condition, the mole fractions of the vapor (liquid) components should be very close to the overall mole fractions $z_i$, and the mole fractions of the liquid (vapor) components approach to zero. On the other hand, if the mixture has two phases at the given condition, the phase identification can be specified by comparing the phase density or phase molecular weight to that of the feed composition. The heavier phase is the liquid phase while the lighter phase is the vapor phase. It can be seen that the predictions of the deep learning model near the phase boundary agree with iterative flash solutions very well in these two cases.
3.3. Prediction Evaluation

The accuracy of our deep learning model shows as the small prediction errors in Table 1. Here we visually compare molar compositions computed by the deep learning and iterative flash approaches to further illustrate this accuracy. Figure 6 displays the mole fractions of C$_1$, C$_{5-7}$ and C$_{10+}$ in the vapor phase of the Bakken oil as a function of the temperature at $C' = 343$ mol/m$^3$. The open and solid symbol represent the deep learning estimation and iterative flash solution, respectively. Even though the mole fraction profiles computed
by both approaches exhibit some opposite variation tendency, the deep neural network is considered to yield good predictions under certain confidence, especially for the heaviest component \(C_{10+}\). Figure 7 shows the mole fractions of \(C_2\), \(C_{5-6}\), \(C_{7+}\) and \(C_{13+}\) remain constant at the specified temperature interval, implying the EagleFord1 oil is in the single-phase state at the given overall concentration. It can be seen the deep learning predictions agree with the iterative flash solutions very well. Figure 8 presents the mole fractions of \(C_1\), \(C_3\), \(nC_4\) and \(CO_2\) in the liquid phase of the EagleFord1 oil at another temperature interval when \(C = 1810\) mol/m\(^3\). The mole fraction of \(C_1\) predicted by the neural network model deviates from the results of NVT flash calculation more distinctly than the other light components. Overall, the optimized deep neural network exhibits a good performance on the prediction accuracy. Admittedly, in some cases there are some slight differences.
Figure 6: Mole fractions of $C_1$, $C_{5-7}$ and $C_{10+}$ in the vapor phase of Bakken oil predicted by the deep learning approach (open symbols) and iterative flash calculation (solid symbols).
Figure 7: Mole fractions of C$_2$, C$_{5−6}$, C$_{7+}$ and C$_{13+}$ in the single-phase EagleFord1 oil. The open and solid symbols represent the deep learning prediction and iterative flash solution, respectively.
Figure 8: Mole fractions of $\text{C}_1$, $\text{C}_3$, nC$_4$ and CO$_2$ in the liquid phase of the EagleFord1 oil. The open and solid symbols represent the deep learning prediction and iterative flash solution, respectively.

The effect of data size on the prediction accuracy is validated in Figure 9. With the temperature increasing, the mole fraction of $\text{C}_1$ keeps constant at its overall mole fraction 0.5816, denoted by the black line, which means the EagleFord1 oil is single phase under the given conditions. As expected, the less data we use to train the model, the more deviation takes place in the outcomes of the deep neural network. It is found that the deep neural network with the data size of $201 \times 201$ gives pretty good approximations, which exhibit some slight fluctuations in comparison to the predictions based on the $301 \times 301$ data size. When the data size increases from $201 \times 201$ to $301 \times 301$, the insignificant accuracy improvement, shown in Figure 9, is not worthy of doubling the computational time as shown in Table 1.
Thus, a moderate number of data should be used to train the model in order to balance the accuracy and efficiency of the deep learning model.

Figure 9: Mole fraction of C\textsubscript{1} predicted by the deep learning model under different data size when the EagleFord1 oil is at the single-phase state. The black line represents the solutions of iterative flash calculation.

4. Conclusions

In this study, we develop a deep neural network to approximate NVT flash calculation so as to speed up phase equilibrium calculations. A dynamic model, comprising the mole and volume evolution equations, is iteratively solved to generate training data for the proposed neural network model. Three real reservoir fluids, including a five-component Bakken oil and
two Eagle Ford oils with eight and fourteen components, are investigated, showing that both
the dynamic flash model and deep learning model can handle complex fluid mixtures. To
increase the prediction accuracy, we reformulate the loss function and employ the dropout
technique to reduce the overfitting problem inherent in neural networks. Furthermore, batch
normalization is applied to accelerate the training process of the deep learning model. It is
noted that the network configuration is designed based on our previous research, which stud-
ied various effects on the performance of deep neural networks for phase equilibria problems.
Different from the preceding research by machine learning models, our deep learning model
gets rid of the constraint of the conventional flash framework where the stability test precedes
phase split calculation, so that it can determine the number of phases without additional
stability testing. In addition, the proposed network model is able to identify the single-
phase state, either single vapor or liquid. A variety of examples are presented to exhibit
the accuracy and efficiency of the proposed neural network model. We find that training a
deep learning model becomes much more expensive above a threshold data volume, however,
with less enhanced accuracy. In our tests, the trained model yields much faster predictions,
at most 244 times for the investigated cases than the iterative NVT flash, and meanwhile
preserve good accuracy. It is worth mentioning that this study paves the path for our future
research on acceleration of phase equilibrium calculations. In addition to further improving
the accuracy and robustness, the proposed neural network model will be extended to varying
composition cases that often take place in EOR processes. Some important effects in
unconventional reservoirs, including capillary pressure and nanopore confinement, will also
be taken into account in the future work to validate the capability of deep neural networks
in confined phase equilibrium problems.
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Supporting Information Available

Molar compositions and compositional parameters of the Bakken, EagleFord1 and EagleFord2 oils.

This material is available free of charge via the Internet at http://pubs.acs.org/.

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