paper 8

Construction and Building Materials 336 (2022) 127557

Contents lists available at **ScienceDirect**

Construction and Building Materials

journal homepage: www.elsevier.com/locate/conbuildmat

Predicting compressive strength of alkaliactivated systems based on the network topology and phase assemblages using tree-structure computing algorithms

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Keywords:

Alkaliactivated system Compressive strength Topological constrainttheory Thermodynamic simulation Machine learning A B S T R A C T

Alkaliactivated system is an environment-friendly, sustainable construction material utilized to replace ordinary Portlandcement(OPC)thatcontributesto9%of theglobal carbonfootprint [Code a1]. Moreover, the alkali-activated system has exhibited superior strength at early ages and better corrosion resistance compared to OPC. [Code a1] The currentstateofanalytical andmachine learningmodelscannotproducehighly reliable predictions of the compressivestrengthofalkaliactivatedsystemsmadefromdifferenttypesofaluminosilicate-richprecursors owing to substantive variation in the chemical compositions and reactivity of these precursors [Code a2]. In this study, a random forest model with two constraints (i.e., topological network and thermodynamic constraints) is employedtopredictthecompressivestrengthofalkaliactivatedsystemsmadefrom26aluminosilicate-rich precursorsanddistinct processing parameters [Code a3]. Results show that once the model is rigorouslytrainedand optimized, the RF model can yielda priori, highfidelity predictions of the compressive strength in relation to the physicochemicalpropertiesofaluminosilicate-richprecursors; processingparameters; and constraints [Code a4]. The topological network constraint provides the chemostructural properties and reactivity of the aluminosilicate-rich precursors [Code a4]. Whereas the thermodynamic constraint estimates the phase assemblages at different degrees of reaction of the aluminosilicate-rich precursors [Code a4]. Finally, the correlations between topological network constraint; phase assemblage; and compressive strength are demonstrated. [Code a5] When the topological network constraint equals 3.4, the alkaliactivated systems can achieve their optimal compressive strength. [Code a5]

1. Introduction

OrdinaryPortlandcement(OPC)isafundamentalmaterialfor infrastructure development. The global production of OPC is 4 billion tons per year with a growth rate of 80 million tons per year[1]. This giganticproduction presentsatremendous energyconsumption(11 exajoule per year[2]) and 9% of the global CO₂emission[3]. Calcination of limestone above 1450 °C contributes at least 50% of CO₂ release in OPC manufacture, where sustainability cannot be improved through clean energy [4]. Another shortcoming faced by OPC is susceptibility todegradation, which results in additional OPC consumption in the form of

repairandreconstruction. With the increasing demand of infrastructures, an alternative cementitious material—more sustainable and durable than OPC—is urgently needed. Alkali-activated system (including binder; mortar; and concrete)—also named as geopolymer system—is a rising candidate to replace OPC [5,6]. The alkali-activated system can be made from aluminosilicate-rich precursors such as fly ash; ground granulated blast-furnaces lag; rice huskash; and other urban waste ashes [6,7]. Although they are made from recycled materials, the alkali-activated system sexhibits trongers trength at early ages and better corrosion resistance than OPC [7–9]. The replacement of OPC by the alkaliactivated system would reduce 73% of CO₂ emission and 43%

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https://doi.org/10.1016/j.conbuildmat.2022.127557

Received 21 February 2022; Received in revised form 31 March 2022; Accepted 15 April 2022

Available online 23 April 2022

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of energy consumption in the cement and construction industry[10]. Alkaliactivated systems are made from amorphous aluminosilicate-rich precursors and alkali activators (e.g., sodium silicate; sodium hy-droxide; potassium silicate; etc.) through a geopolymerizationreaction[5,8]. The alkaliactivator significantly increases the c oncentration of cations and OH⁻ in the solution, resulting in an acceleration of leaching of aluminate and silicate ions from surfaces of aluminosilicate-rich precursors. Afterthis, sodium aluminosilicate hydrate (N-A-S-H) and calciumsodiumaluminosilicate hydrate(C-N-A-S-H)gelsareprecipi-tated as a result of the geopolymerization reaction. (N.B.: N=Na₂O; A=Al₂O₃; C = CaO; $S = SiO_2$; $H = H_2O$). The dominant product of Ca-rich precursorisC-N-A-S-H; otherwise, the dominant product is N-A-S-H. Conventional (twopart) alkaliactivated systems are produced by mix-ingaluminosilicaterichprecursors; alkaliactivators olutions; and additional water [11]. Due to the danger of handling large amounts of theviscous and corrosive alkaliactivator solutions, one-partalkali- activated systems have been widely researched upon. One-part alkaliactivated systems are synthesized by the addition of water to dry pre-mixture, prepared from aluminosilicate-rich precursors and solid alkaliactivators with or without calcination[8].

Reactivity of aluminosilicate-rich precursors is a vital parameter that affectsfreshandhardenedpropertiesofalkali-activatedsystems[12,13]. The highly reactive aluminosilicate-rich precursors setina short time and yield high compressive strength. The reactivity of the aluminosilicate-rich precursors is associated with chemostructural

properties of precursors, especially the amorphous phases accounting for almost 50–90% in aluminosilicate-rich precursors [14,15]. Previous studies [15,16] have shown that the topological constraint theory (TCT) is the keyenable rto evaluate the reactivity of aluminosilicate-rich precursors. The TCT is developed based on the topology of the atomic structure of amorphous materials and their macroscopic properties [17]. To be specific, TCT reduces the complexity of chemostructural properties of amorphous materials into a singular constraint [number of constraints (n_c)] [18]. This parameter quantitatively evaluates the reactivity of aluminosilicate-rich precursors by solely relying on the major components (i.e., CaO; SiO₂; and Al₂O₃).

Compressivestrengthisoneofthemostimportantmechanical properties of alkaliactivated systems, giving a general idea about the performance regarding the quality of alkaliactivated systems[19,20]. Foralkaliactivated systems to achieve the targeted compressive strength as well as other mechanical properties, researchers must invest a significant amount of time in exploring mixture design and processing parameters [21]. The laboratory exploring work, however, is costly and labor-intensive. Thus, being able to predict the compressive strength is essentialespeciallyinsavingtimeandcost.Consequently,manyresearchershaveworkedonvariousanalyticalmodelstopredictthe compressivestrengthofalkali-activatedsystems.Leetal.[22] have developedamodifiedFeret'smodelbasedonthemixturedesignof alkaliactivated systems and standard compressive strength at 28-day. Thismodel, however, cannot properly predict the compressive strength of the systems without aggregate [23]. An analytical model forcompressive strength of alkaliactivated systems made from a fly ash has been developed by Beluah et al. [24]. However, the shortcoming is that a calibration is required when the model is applied to a new database due to alimited number of data-records used for the model development. JonbiandFulazzaky[25] haverelated compressivestrength with the ageofalkali-activatedsystems. This study cannot produce reliable predictions ofcompressive because of exclusion of some influential variables. This study also reinforces that analytical models cannot account for all influential

variables and variables without understanding their contributions on

properties.

Machine learning (ML), a data-driven artificial intelligence, can overcometheabovementionedshortcomingsofcurrentanalytical models.Inthepastdecades,manystudies[19,26–30] haveapplied machinelearningmodelstopredictdifferentpropertiesofalkali-activated systems. Zhang et al. [29] have shown regression-and tree-basedMLmodelstoproducereliablepredictionsofcompressive strengthofalkali-activatedsystemsinrelationtomixturedesignand chemistryof precursors.Gomma etal.[28] havedemonstrated the reliable performance of random forest (RF) model on predicting fresh and hardened properties of alkaliactivated systems related to mixture design and various processing conditions. Lahoti et al.[30] have used RF, Naïve Bayes, and k-nearest neighbor models to predict mechanical properties of alkaliactivated systems with a small database.Although previous studies have shown that machine learning is a promising tool to predict properties of alkaliactivated systems, some knowledge gaps still

exist.First,previousstudiesonlyincludeoneortwotypesof aluminosilicate-rich precursors in their databases. The performance of ML models on multiple (more than two) aluminosilicate-rich precursors

hasnotbeenvalid.Next,asaforesaid,thereactivityofthe aluminosilicate-rich precursors is a critical parameter that influences the properties of the alkaliactivated systems. The parameter related to the reactivity has not been included in previous studies. Finally, ML models find the underlying structure between the mixture design and

properties, all without understanding mechanisms. This leads to a critical problem that predictions as produced by ML may violate fundamental material laws. Such violation significantly undermines the credibility of results from ML. However, this problem can be solved by applying thermodynamic constraints to ML models. Han et al. [31] have shown that the thermodynamic constraint regulates the RF model to predict properties of OPC systems, where the accuracy is better than models without constraints.

Gibbs Energy Minimization Software (GEMS) is a tool for geochemicalmodeling[32,33],whichcanbeemployedtoconduct thermodynamicsimulationsofalkali-activatedsystems. Morespecifically, GEMS simulations and the degree of reaction of the aluminosilicate-rich precursors can produce equilibrium phase as semblages and

microstructure for alkaliactivated systems. Several studies[34,35] have shown that the phase assemblages from GEMS simulations accurately replicate phases appearing in real alkaliactivated systems. In thethermodynamicsimulations, the expected products for alkaliactivated systems are: C-N-A-S-H; N-A-S-H; stratlingite; gibbsite; monocarboaluminate hydrate; hydrogarnet; and amorphous zeolites. With the understanding of phase assemblages at different degrees of reaction of aluminosilicate-rich precursors, microstructures and properties of the alkaliactivated systems can be correlated. This would help researchers understand the mechanisms behind properties of alkaliactivated systems.

The study presented here aims to overcome the above mentioned limitations of current analytical and ML models by advancing network andthermodynamicconstraintsthatenhancethepredictionperformanceofMLmodelsonthecompressivestrengthofalkali-activated systems. Herein, the RF model is employed to find correlations betweencompressivestrengthandinputs(i.e.,mixturedesignandprocessingparametersofalkali-activatedsystems). Such correlations are utilized to produce a priori, reliable predictions of compressive strength innewalkaliactivatedsystems. The network constraint (developed from chemostructural properties of a luminosilicate-rich precursors basedonthetopologicalconstrainttheory) and thermodynamic constraint (obtained from thermodynamic simulations) are employed to regulatetheRFmodel, which ensures that predictions do not violate fundamentalmateriallaws.Furthermore,theRFmodellinksthe compressivestrengthwithphaseassemblagesofalkali-activatedsys- tems. This leads to an investigation of compressive strength behavior in relation to microstructures reacted from aluminosilicate-rich precursors with different chemostructural properties.

1. **Modeling** method

Inthisstudy,aRFmodelisemployedtopredictthecompressive strengthofalkaliactivated systems.First, themodelis trainedand tested without any number of constraints; described inSection 2.1) is applied to the model toproduce more accurate predictions. Third, an additional thermodynamic constraint (volume fraction of products; described in Section 2.2) regulates the networkconstrained model to ensure non-violation offundamentalmateriallaws. Prediction performances of three stages are rigorously appraised by comparing their prediction against measured values. Fig. 1 demonstrates the framework of the RF model with constraints. As can be seen, the model used in this study unites the network and thermodynamic constraints with the RF model to achieve optimal predictions of compressive strength of alkali-activated systems. The purpose of using three parallel predictions is to investigate the optimal prediction strategy (one that treads the balance between ease-of-use/simplicity and accuracy) for alkali-activated systems. The optimal strategy can be directly applied to the database without other two stages. A detailed description of the RF model can be found in Section 2.3.

2.1. Network constraint

Herein, a brief description of TCT is given below, and details can be found elsewhere [16]. The major components (i.e., SiO_2 , CaO, and Al_2O_3) of the aluminosilicate-rich precursors are utilized to derive thenumber of constraints based on the topological constraint theory (TCT). The compositions of the aluminosilicate-rich precursors are considered as $(CaO)_x(Al_2O_3)_y(SiO_2)_{1-x-y}$, wherexandyare normalized molar fractions. Aluminosilicate-rich precursors contain radial bonding-stretching (BS)andangularbond-bending (BB)constraints[16,18,36].BSexists between two bonded atoms. BB exists when atoms have fixed interatomicangles. There are sixtypes of atoms that have constraints: Si atom; Al atom (four-fold and five-fold); tricluster oxygen atom (TO); free oxygen atoms (FO); bridging atom (BO); and non-bridging atom (NBO). In the

aluminosilicate-rich precursors, Si and Al atoms are the networkformers, forming tetrahedral structures. Oatoms acting as a bridge to connect Si/Al tetrahedrons are BOs [37,38]. The Ca atom is the network modifier existing between Si/Al tetrahedral molecular as the interstitial site[37,38]. The O atom connected with Ca atom and Si/Al tetrahedral molecular is NBO. FO is an O atom that only connects with a Ca atom. TO is the O atom acting as the charge compensating atom for theAltetrahedron.DependingontheCaOandAl₂O₃contents,the network structure of aluminosilicate materials can be divided into three regimes: fully depolymerized (y —x \leq —²); partially depolymerized (—² \leq y —x \leq 0); and fully polymerized $(0 \le y - x)$ [16]. In the fully depolymerized regime, Ca atom is the dominant composition. All Si and Al tetrahedrons isolate from each other. The excess Ca atoms result in the formation of NBOs and FOs. In the partially depolymerized regime, Si atom is the dominant composition. The Ca atoms connect with Si/Al tetrahedronsandcompensatechargeofAltetrahedrons. This regime includes both BOs and NBOs. In fully polymerized regime, Al atom is the dominant composition. Due to insufficient Ca atoms, the charge of Al atoms cannot be compensated to form four-fold tetrahedral units, where over-coordinated Al atoms (five-fold) are formed. This regime includes both BOs and TOs.

Caatomisthecharge-compensatingion; thus, it is excluded from topological constraints. But it creates 1 BS constrain with a connected NBO[39]. Ca atom connected to a FO can form 1 BS constraint. Previous studies [16,36,39–41] have shown that Siatoms form tetrahedral structure at five fixed angles in the amorphous phases, which contains 4 BS and 5BB constraints. Additionally, Sitetrahedron creates 1BB constraint with BO. Four-fold Alatomhas 4BS constraints and 5BB constraints. Five-fold atom only has 5BS constraints. To has 3BB constraints. The number of constraints (n_c) corresponding to each regime is shown in **Eq.1**. Additional emphasis is given to that the aluminosilicate rich precursors in this study fall into partially depolymerized and fully polymerized regimes.

3

$$n_c$$
 = 11 + y — 10x (Fully depolymerized) (1a) x y
3 — 2 + 2
 n_c = 11 + 10y — 10x (Partially depolymerized) (1b) x y
3 — 2 + 2
 n_c = 11 + 13y — 13x (Fully polymerized) (1c) x y
3 — 2 + 2

2.2. Thermodynamic modeling

Fig. 1. Schematic of the RF model to predict the compressive strength of alkali-

activated systems by incorporating network and thermodynamic constraints. Network constraint is developed from the topological constraint theory. Ther-modynamic constraint is acquired from phase as semblages of alkaliactivated systems.

GEMS[32,33] is employed to produce thermodynamicsimulations andphaseassemblagesofalkali-activated systems in relation to their mixture design [i.e., chemical composition of binders (aluminosilicate-rich precursors; alkaliactivators; and water)] and processing conditions (i.e., temperature). The quality of the thermodynamic simulation results is dependent upon the accuracy and completeness of the input properties of the substances and phases, which can typically be found within literature and thermodynamic databases. The thermodynamic data for aqueous species

and numerous solids are acquired from the PSI-GEMS thermodynamic database, while solubility products for relevant phases are extracted from the Cemdata 18 [42,43] and zeolite 20 [44]. An extended Debye-Huckel calculation is utilized by the software to estimate the activity coefficients of the aqueous species. The assumption that the aqueous phase is dominated by NaOH is made. The average ion size and parameter for common short-range interactions of charged are

1. 31Aand0.098kg/mol,respectively.Allphaseassemblagesare simulated at 1 atm and 20-to-85°C.

Phaseassemblagesobtainedfromthethermodynamicsimulations are shown in Fig. 2. The figure reveals volume of all reactants and products a tincreasing degrees of reaction of the aluminosilicate-rich precursors. In Fig. 2, it has been observed that GEMS can estimate phase as semblages of alkaliactivated systems made from multiple aluminosilicate-rich precursors at different water-to-solid ratio. The phases shown herein are in agreement with previous studies [20,34,45]. The main phases of alkaliactivated systems include: C-N-A-S-H; N-A-S-

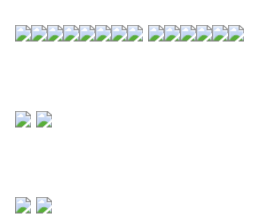


Fig. 2. Equilibrium phase assemblage, simulated using GEMS, of alkaliactivated systems made from (a) fly ash and ground granulated blast-furnace slag (GGBS); and(b)redmudandgroundgranulatedblast-furnaceslag(GGBS). The vertical dashed line presents the phase assemblage at the

targeteddegreeofreaction of aluminosilicate-rich precursors estimated from the compressive strength. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

H (Na-chabazite; zeolite 4A; hydroxysodalite; and natrolite); amorphous Si; Ca-chabazite; Gibbsite;straetlingite; and C₃AH₆. Thephase assemblage is an important outcome because researchers can use it to estimate thevolumefractionofproductsandporosityinanygiven aluminosilicate-richprecursoratthetargeteddegreeofreaction. The compressive strength can be roughly estimated from porosity and volumefractionofproductsbecausetheyprovideabasicsolid-to-solid connectivitywithinthealkali-activatedsystems[46–48]. Inorderto obtain accurate phase assemblages, it is essential to identify the degree ofreactionofthealuminosilicate-richprecursors, which can be estimated directly from compressive strength at any given age. The method to determine the degree of reaction is explained in **Section 3.0**. The volume fraction of products is employed as the thermodynamic constraint to regulate the RF model.

2.3. Random forest (RF) model

RF model is a tree-structure model developed from the classification-and-regressiontrees(CARTs)modelwithbaggingtechnique[49,50]. During the training, the model parallelly constructs hundreds of inde-pendent CARTs from bootstraps. Each tree grows through binary splits in a recursive fashion until the terminal nodes reach a "near-homoge- nous" state. The model allows each CART to grow to its maximum size withoutpruningandsmoothing. WhennewdataisappliedtotheRF model, the model averages the predictions from all trees to produce the final output. Compared to other ML models, the RF model is different in the sense that two-stage randomization[50,51] is employed during the growth of CARTs. The first randomization is that the bootstrap used to constructatreerandomlyselectsdatafromtheoriginaldataset. The secondstage involves that arandomly chosen subset of variables, instead of

using all variables, is utilized to ascertain the best scenario for each split. Due to the two-stagerandomization, trees in the forest decorrelate with each other, resulting in a reduction of variance errors [52,53]. To achieve the RF model's best performance, 10-fold cross-validation (CV) method [54,55] and the grid-search method [28,56] are employed to determine the optimal hyperparameters for the RF model. In this study, 300 trees and 4 splits at each node are for all predictions.

To train and validate the RF model, the database (shown inSection **3.0**)issplitintotwonon-overlappingdatasets:trainingdatasetandtesting dataset. The training dataset contain 75% of randomly selected data-recordsformtheparentdatabase,andtheremaining25%data-recordsformthetestingdataset.Thetrainingdatasetisusedtotrain theRFmodel rigorously.Thetestingdatasetisusedtoquantitatively evaluate the prediction performance of the RF model through 5 statis-ticalparameters—Pearson correlation coefficient (R); coefficient of determination (R²); mean absolute percentage error (MAPE); root mean squarederror(RMSE);andmeanabsoluteerror(MAE).Mathematical functions for these statistical parameters were detailed in our previous studies[54,57,58].

2.4. Database Collection

Data pertaining to compressive strength in 520 alkaliactivated systems[includingpurebinder(aluminosilicate-richprecursors+alkali activator+water); mortar(binder+fine aggregate); and concrete (binder+ fine aggregate+ coarse aggregate)] were collected from 15 studies[59–73]. Majority of the alkaliactivated systems were synthesized byfly ash andground granulated blast furnaceslag, andthe remainingsystemsweresynthesizedfromflyashsinkingspherical beads, red mud, metakaolin, silica fume, and rice husk ash. An alkaliactivatedsystemcancontainatthemostthreedifferent aluminosilicate-richprecursors. Notallthedensitiesofthe aluminosilicate-rich precursors were illustrated in previous studies. In ourdatabase, densitiesofthealuminosilicate-richprecursorsare

calculated based on the density of CaO(3.34g/cm³); Na₂O(2.27g/cm³); SiO₂ (amorphous, 2.20 g/cm³)[74]; and Al₂O₃ (amorphous, 2.32 g/cm³) [75,76] with respect to their normalized mass fractions. Other compositions (e.g., MgO; K_2O ; etc.) are minor, and their products have negligible contribution to the compressive strength [7]. Therefore, they are excluded from the density calculations and thermodynamics imulations. It is worth noting that this database includes not only two-part (aluminosilicate-rich precursors mixed with an alkaliactivator solution) but also one-part (aluminosilicate-rich precursors and solid alkaliactivator mixed with water) alkaliactivated systems.

Here, compressive strength is used as a direct indicator to calculate the degree of reaction of the aluminosilicate-rich precursors. Previous studies[77,78] havedemonstrated that compressive strength of cementitious materials and the degree of reaction of binder exhibits a linear correlation. The degree of reaction of the aluminosilicate-rich precursors is employed to locate the phase assemblage of the binder in thermodynamic simulations. The volume fraction of products obtained

fromthephaseassemblagesactasathermodynamicconstraintto regulate and improve predictions as produced by the RF model. A pre-processing is required before the degree of reaction is correlated with the compressive strength. In this database, the compressive strength was testedfrombinder;mortar;andconcrete. Fuetal.[79] havedemonstratedthatcompressivestrengthisimprovedbyincreasingbinder content and decreasing aggregate content. To unify the degree of reactionforallspecimensinourdatabase, it is important to normalize measured compressive strength by a standard specimen composition (i. e.,astandardbinder-to-aggregateratioinvolume). This is because inconsistent binder-to-aggregate ratios result in inaccurate correlations betweencompressivestrengthandthedegreesofreactionofthe aluminosilicaterichprecursors. The compressive strength of pure binder systems is directly used to determine the degree of reaction. For theremainingalkali-activated systems, the volumeratio of binder-toaggregateiscalculated, and most systems fall into the 0.4-to-0.6

range. Thus, 0.5 of binder-to-aggregateratio is selected as the standardcompositiontonormalizecompressivestrength.Here,the normalized compressive strength ($CS_{normalized}$) is calculated on the basis of **Eq.** (2). CS_{measured} is compressive strength measured from experiment;r_{measured} is the volume ratio of binder-to-aggregate of each specimen; andr_{standard}isthevolumeratioofbinder-to-aggregateofthestandard specimen, which is 0.5. The reactivity (maximum degree of reaction) of the aluminosilicate-rich precursors depends on the crystallinity, where a highlycrystalline aluminosilicate-rich precursors results in a low reactivityandthusyieldingalowcompressivestrengthinthealkali- activated systems[12]. Previous studies[14,15,80] have revealed that theamorphous content of the aluminosilicate-rich precursors ranges from 40% to 90%. This indicates that the maximum degree of reaction of the aluminosilicate-rich precursors cannot exceed 90%. In our study, we assume that 80% of the aluminosilicate-rich precursors have reacted in the specimen with the highest compressive strength (80 MPa). The de-gree of reaction of the aluminosilicate-rich precursors for the remaining specimens is computed as the proportion of 80% based on the fraction of the compressive strength to maximum compressive strength.

Table 1
Statistical parameters pertaining to 10 inputs; 2 constraints; and 1 output (bold) of 520 alkali activated systems in the compressive strength database.

Attribute	Unit	Min.	Max.	Mean Std. Dev.
Aluminosilicate Material Content	%mass	9.451	73.76	26.25 16.64
Fine Aggregate Content	%mass	0	66.67	32.33 16.99

Coarse Aggregate Content	%mass	0	79.55	27.33	3 23.07
Na ₂ SiO ₃ Content	%mass	0	17.243	4.420	3.600
NaOH Content	%mass	0	3.194	0.857	7 0.821
Superplasticizer Content	%mass	0	0.427	0.075	5 0.141
Water Content	%mass	2.685	5 25.92	8.395	5 4.073
CuringTemperature	۰C	20	85	36.47	7 22.94
Curingtime	Days	1	28	12.84	10.75
Type	unitless	s 1	2	_	_
Number of Constraints	unitless	s 3.092	2 4.155	3.634	1 0.297
Volume Fraction of Product	unitless	s 0.02 3	0.6921	0.214	1 0.136
Compressive Strength	MPa	5.152	79.82 27.16 16.80		

CSnormalized = CSmeasured rmeasured

rstandard

(2)

Fig. 3. A linear correlation between compressive strength and volume fraction

The database used for ML models contains 10 inputs and 1 output. The inputs include: normalized mass of aluminosilicate material; solid NaOH;solidNa₂SiO₃;fineaggregate;coarseaggregate;andwater (%_{mass}); curing temperature ($^{\circ}$ C); age (days); type (unitless; 1 = one-part and 2= twopart). The output is compressive strength (MPa), which is the measured value obtained from the literature. It is worth noting that NaOH and Na₂SiO₃are alkali activators. Thenumber of constraintsof thealuminosilicate-rich precursors (unitless) and volume fraction of products (unitless) are applied to the model as additional inputs when the network and thermodynamic constraints are applied. It should be noted thatthenumberofconstraintsofthealuminosilicate-richprecursorsis calculated by adding up the number of constraints of each aluminosilicaterichprecursorwithrespecttoitsmassfractioninall aluminosilicate-rich precursors. Statistical parameters pertaining to in-puts and output are itemized in Table 1. The database used in this study is provided inSupplementary Information.

1. **Results and** discussion

3.1. Thermodynamic simulations

Based on GEMS simulations, we find that across all alkaliactivated systems resulted in a near-linear correlation (Fig. 3) with the compressivestrengthagainstthevolumefractionofproducts. Thereason to evaluate this relationship is that the compressive strength of any alkaliactivated systemis correlated with the extent of reaction of the aluminosilicaterich precursors, which, in turn, dictates the volume of products in 520 alkaliactivated systems used in this study. A mathematical function qualitatively estimating this correlation is shown in the legend.

fraction of the products. It should be noted that a mathematical function thatinterpretsthecorrelationbetweencompressivestrengthandvol-ume fraction of products in 520 alkaliactivated systems is shown in the legend. This mathematical functional lows scientists to promptly examine the compressive strength of any given alkaliactivated system by using its volume fraction of products as the sole input.

ItisworthpointingoutthattheRFmodelcandirectlyproduce predictionsofthevolumefractionofproductsinalkali-activatedsys- tems, without calculating them based on phase assemblages from thermodynamic simulations. This requires the RF model to be trained with a new output, which consists of the same input variables shown in **Section**3.0,butthevolumeofproductsastheoutput.OncetheRFmodelis trained, it can predict the volume fraction of products (Fig. 4) in new alkaliactivated systems with respect to their mixture design. The predicted volume fraction of products can also be used as the thermodynamic constraint(the same constraintobtainedfrom thephase assemblage)toregulatetheRFmodeltopredictthecompressive strength.However,inthisstudy,allthermodynamicconstraintsare derived from the GEMS simulations.

3.2. Machine learning predictions

Fig.5 demonstrates the predictions of the compressive strength of alkaliactivated systems produced by the RF model without constraints



Fig. 4. Predictions of volume fraction of products as produced by the RF model against values obtained from thermodynamic simulations. Pearson correlation

coefficient(R)ofpredictedresultsisshowninthelegend. The dashline indicates ideal prediction, and solid lines represent a±10% error bound.



Fig. 5. Predictions of compressive strength of alkali activated systems as pro—ducedbytheRFmodelwithoutanyconstraintscomparedagainstmeasured compressive strength. Pearson correlation coefficient (R) of predicted results is

showninthelegend. The dash line indicates ideal prediction, and so lid lines represent a $\pm 10\%$ error bound.

against the measured compressive strength. Predictions for both training andtestingdatasetsareshowninthefigure. The five statistical parameters (mentioned in Section 2.3) pertaining to the performance of the RF model on the testing dataset are itemized in Table 2.

As shown in Fig. 5 and Table 2, the accuracy of predictions produced bytheRFmodelismoderate.R² is0.63andRMSEis9.752MPa. Although

previous studies [19,26,28,81,82] have demonstrated that

Table 2

The statistical parameters pertaining the performance of the RF model with or withoutnetworkandthermodynamicconstraintsonpredictionsofalkaliactivated systems' compressive strength against the testing dataset.

Model Name	R	R2	MAE	MAPE	RMSE
	Unitless	Unitless	MPa	%	MPa
RF	0.8095	0.6553	6.854	29.39	9.433
RF + Network Constraint	0.8841	0.7816	5.956	24.56	7.868
RF + Network Constraint+	0.9619	0.9253	3.649	13.45	4.865

Thermodynamic Constraint

standaloneMLmodelscanyieldaccuratepredictionsofcompressive strengthofalkali-activatedsystems, it is expected that a standalone model cannot produce highly accurate predictions for the database used in this study. This is because our database is much more complex than databases used in previous studies [19,26,28,81,82]. First, both one-part and two-part alkaliactivated systems are included in the database. In general, due to heat generated from the dissolution of solid activators, one-part alkaliactivated systems are more reactive than two-part systems[83]. Therefore, one-partalkali-activated systems demonstrate highercompressive strengthatearly ages and as horters etting time compared to

two-part systems. Second, the chemical composition of the aluminosilicaterichprecursorsinpreviousstudiesweresimple, only containing a few different compositions. However, there are 26 different compositions of the aluminosilicate-rich precursors in this study. Literature[9,20,84] has shown that chemical compositions of the aluminosilicaterichprecursorssignificantlyaffectsthecompressivestrength of alkaliactivated systems, where high Al₂O₃and SiO₂contentslead to binders with high compressive strength. Third, the database in this study contains a largescale of water-to-solid ratio from 0.18-to-0.50. The geopolymerization reaction is incomplete without sufficient water; in contrast, excessive watercreates an additional porositythatcan reduce compressive strength. Finally, the database consists of a largerangeofcuring temperatures, from 20to-85°C.Elevatingthecuring temperatureincreasestheextentandrateofthereactionthrough escalatingmesoporevolumeandsurfaceareafornucleiofgeo-polymerization reaction [85]. This accelerates setting time accompanied by higher compressive strength at early ages.

As abovementioned, due to the complexity of the database used in thisstudy, it is important to further calibrate the RF model to obtain better predictions of compressive strength. Towards this, the topological network constraint (described in Section 2.1) and volume fraction of products from the rmodynamic simulations (described in Section 2.2) are used to provide information of fundamental material laws to guide the RF model. Fig. 6a demonstrates the predictions of compressive strength of alkaliactivated systems produced by the RF model with the network constraint. Fig. 6b shows the predictions of compressive strength of alkaliactivated systems produced by the RF model with the network and thermodynamic constraints. Predictions for both training and testing datasets are summarized in Table 2.

As can be seen in Fig. 6 and Table 2, the RF model produces accurate predictionsforalkali-activatedsystems' compressive strength with network and thermodynamic constraints. R² and RMSE of the predictions with the network constraint are 0.78 and 7.868 MPa, respectively. R² and RMSE of the predictions with the network and

thermodynamic constraints are 0.93 and 4.865 MPa, respectively. The prediction error is even smaller than the standard deviation (5 MPa) of the compressive strength measurement [86]. Based on these values, it is clearthatthepredictions with constraints are superior compared to predictions produced by the standalone RF model, which is expected. Unlike the standalone model without knowing the chemical composition of the aluminosilicate-rich precursors, the network constraint provides chemostructural properties (e.g., the quantities of NBOs; BOs; FOs; and TOs and strength of chemical bonds) of aluminosilicate-rich precursors to guide the RF model. This information can serve as the proxy for the reactivity of a luminosilicate-rich precursors. The highly reactive aluminosilicate-rich precursors react fast with the alkali activators and waterresultinginhighercompressivestrengthatearlyagesandan overallhigherdegreeofreaction. Thus, critical information from the network constraint helps the RF model to catch the trend between inputs and compressive strength. The thermodynamic constraint refines pre-dictions of compressive strength further. As shown in Fig. 3, compressivestrengthexhibitsastrongcorrelationwiththethermodynamic constraint (volume fraction of products). Therefore, the thermodynamic constraint restricts predictions produced by the RF model in a small and reliable range. Furthermore, the thermodynamic constraint prohibits



Fig. 6. Predictions of compressive strength of alkali activated systems as produced by the RF model with (a) network constraint and (b) network + thermodynamic constraints compared against measured compressive strength. Pearson correlation coefficient (R) of predicted results is shown in the legend. The dash line indicates ideal prediction and solid lines represent a±10% error bound.

the RF model from violating fundamental material laws.

3.3. The role of chemical compositions of aluminosilicate materials

Previous sections have shown that the RF model with constraints can produce reliable predictions of compressive strength. However, because machinelearningmodelsarea"black-box",theycannotinterpret mechanismsbehindcompressivestrengthandmixturedesign. This sectionrevealstheinfluenceofthechemicalcompositionofthe aluminosilicaterichprecursorsandthemicrostructureofalkaliactivated systems on the compressive strength through a relationship between the number of constraints, phase assemblages, and compressive strength. The simulations are conducted with 0.35 of water-to-solid ratio; 0.14 of alkali-to-fly ash ratio; 0.27 of NaOH-to-NaSiO₂ ratio; and 20° C.Theflyashcompositionis(CaO)_{0.4-x}(Al₂O₃)_x(SiO₂)_{0.6},andthe degreeofreactionis60%. Such parameters are chosen because most datarecordsinthedatabasehavesimilarparameters. The simulated results for fully polymerized and partially depolymerized are shown in Fig. 7. As expected, the main phases are C-N-A-S-H and N-A-S-H (Na-Chabazite) [9,45]. Forboth regimes, atn_c=3.4, the compressive strength has the highest value. This is because the trade-off between C-N-A-S-HandN-A-S-Hphasesreachestheoptimal state. Specifically, before $n_c = 3.4$, alkaliactivated systems do not contain enough network forming atoms to form a high strength gel-like polymeric structure.





Fig. 7. Phase assemblages (bottom) of alkaliactivated systems simulated from thermodynamic modeling and compressive strength (top) predicted from the RF

modelcorrespondingtonetworkstructure(numberofconstraints)of(a)fullypoly merizedflyash;and(b)partiallydepolymerizedflyash.Theparametersfor thermodynamic simulations are shown in the legend.

However, aftern_c=3.4, the reactivity of fly ash diminishes due to excessnetwork former atoms.

Fig. 7a demonstrates the phase assemblage and compressive strengthfor fully polymerized fly ash. The Al_2O_3 varies from 0.2 (n_c =3.23) to $0.4(n_c=4.00)$. Withincreasing Al_2O_3 and decreasing CaO_3 , it can be observed that the volume of C-N-A-S-H decrease monotonically. Atn_c = 3.7, Cachabazitestartsforming.MostoftheNa2Ohasformedstable phases(Na-Chabazite)withAl₂O₃,andinsufficientCaOforcesmore Al₂O₃ togetintoaCarichstructure, where producing $1 \text{mol} \text{Ca-chabazite consumes } 2 \text{ mol of } \text{Al}_2 \text{O}_3$. Later CaO is too low to form a Ca-rich structure, thus gibbsite starts forming. With decreasing volume of C-N-A-S-H, the compressive strength values tend to plummet. This reinforces that C-N-A-S-H is the major component to provide strength in alkali-activatesystems[7,8,20].Cachabaziteandgibbsitebarely contribute to compressive strength. Fig. 7b shows phase assemblage and compressivestrengthforpartiallydepolymerizedflyash.TheAl₂O₃varies from 0.19 (n_c = 3.23) to 0.08 (n_c = 4.00). It is clearly observed that volume of C-N-A-S-H decrease with increasing number of constraints. Compared to fully polymerized fly ash, partially depolymerized fly ash producesmoreC-N-A-S-H.Therefore,thecompressivestrengthof partiallydepolymerizedflyashishigher, eventhoughthereismore porosityinthestructure.Partiallydepolymerizedflyashesalsoformlesser amorphous Si phase because they have sufficient CaO and Al₂O₃toform C-N-A-S-H and N-A-S-H phases.

1. Conclusions

ThispaperemployedtheRFmodeltopredictthecompressive strength of alkaliactivated systems made from seven types (26 different compositions)ofaluminosilicate-richprecursors. Inordertoimprove prediction performances, thermodynamic (volume fraction of products ofalkaliactivatedsystems) and network (number of constraints of the aluminosilicate-rich precursors) constraints were utilized to regulate the RF model. The correlations between the phase assemblages, number of constraints, and compressive strength were revealed and explained. It is worthwhile topoint out that this is the first study that employs ML models with constraints to predict the compressive strength of aluminosilicate-rich precursors and wide processing parameters.

The database for alkaliactivated systems contained 520 unique datarecordswith 10 inputs variables. The parent database was split into trainingandtestingdatasetsto trainthemodel andevaluate the performance. The prediction accuracy (R = 0.81) of RF model without any constraint is acceptable. The prediction accuracy (R=0.88) of RF model with network constraint is moderate. The prediction accuracy (R= 0.96) of RF model with network and thermodynamic constraints is highly reliable. Results sufficiently reveal that the network and thermodynamic constraints provide necessary chemostructural information and material laws to enhance the performance of the RF model. The well-trained RF modelwithconstraintswasusedtoestablishcorrelationsbetween microstructure of alkaliactivated systems; reactivity of aluminosilicaterichprecursors; and compressive strength. When the number of constraints was eq ualto 3.4, the alkali-activated systems achieved the optimal compressive strength. The C-N-A-S-H phase is the major phase that provide this high compressive strength.

Inconclusion,thepredictionsshowninthisstudycanstillbe improved with a larger and more diverse database. In the future, except forAl₂O₃,CaO,SiO₂,otherchemicalcompositionsofthe aluminosilicate-

richprecursorscanbeaccountedinthephaseassem- blage. This is towards understanding the influence of minor components and theirproducts on compressive strength. However, this paper pre-sentsanewpathway tounderstandingmicrostructure-property correlations. Meanwhile, the ML models can provide a preview of properties of alkaliactivated systems made from new aluminosilicate-rich precursors without performing experiments.

CRediT authorship contribution statement

Rohan Bhat: Taihao Han: Sai Akshay Ponduru: Arianit Reka: Conceptualization. JieHuang: GauravSant: Conceptualization. Adity a Kumar: Conceptualization.

Declaration of Competing Interest

Theauthorsdeclarethattheyhaveno knowncompetingfinancial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgement

This study was financially supported by: the Leonard Wood Institute (LWI:W911NF-07-2-0062);theNationalScienceFoundation(NSF-CMMI: 1661609; NSF-CMMI: 1932690; NSF-DMR: 2034856); and the Federal Highway Administration (Award no: 693JJ31950021); and the Fulbright Visiting Scholar Program.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.conbuildmat.2022.127557.

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