A new method to predict coronary artery lesions (CALs) in Kawasaki disease (KD) was developed using a mean structure equation model (SEM) and neural networks (Nnet). There were 314 admitted children with KD who met at least four of the six diagnostic criteria for KD. We defined CALs as the presence of a maximum z score of ≥ 3.0. The SEM using age, sex, intravenous immunoglobulin resistance, number of steroid pulse therapy sessions, C-reactive protein level, and urinary β2-microglobulin (u-β2MG/Cr) values revealed a perfect fit based on the root mean square error of approximation with an $R^2$ value of 1.000 and the excellent discrimination of CALs with a sample score (SS) of 2.0 for a latent variable. The Nnet analysis enabled us to predict CALs with a sensitivity, specificity and c-index of 73%, 99% and 0.86, respectively. This good and simple statistical model that uses common parameters in clinical medicine is useful in deciding the appropriate therapy to prevent CALs in Japanese KD patients.
and without IVMP. This enabled us to develop a new method of predicting CALs early during admission to help us choose the appropriate therapy for KD patients.

We focused on the pathogenesis of KD, which is associated with some gene and protein abnormalities. To analyse the unknown factors, we used a structure equation model (SEM) analysis since we could assume the role of unknown factors in the study using a latent variable. We used a sample score (SS), a factor score for latent variables calculated using the AMOS software, because it is thought to reflect the influence of a latent variable on each subject. We also used a Nnet analysis to predict CAL formation because it is useful for efficiently predicting several events.

Finally, we found the importance of the SS to remarkably discriminate the occurrence of CALs. Thereafter, we introduced an efficient and simple statistical model using the Nnet analysis, which enabled us to predict CALs with sensitivity, specificity and c-index values of 73%, 99% and 0.86, respectively. This method is useful in deciding the second- or third-line therapies using the anti-tumour necrosis factor, cyclosporine, or plasma exchange for Japanese KD patients on admission after first- or second-line therapy using IVIG with steroid hormones.

Results

Amongst the 375 children clinically diagnosed with KD, 61 were excluded. Finally, 314 patients including 106 of 118 in the first study and 208 of 257 in the second one were included (Fig. 1). The patient profiles in the first and second studies are described in Table 1. The value of log urinary β2-microglobulin (u-β2MG/Cr) and the IVIGR value were significantly higher in the first study, whereas the age and coronary artery diameter before the treatment were significantly higher in the second study.

Table 1. Comparison of basic characteristics between the first and second studies. Statistical analysis was performed using Student’s t test and chi-square test. *p < 0.05, **p < 0.01. SD standard deviation, n number, IVIGR intravenous immunoglobulin resistance, IVMP times number of intravenous methylprednisolone pulse treatment courses, WBC white blood cell, CRP C-reactive protein, T-Bill total bilirubin, AST aspartate aminotransferase, u-β2MG urinary β2-microglobulin.
Clinical risk factors associated with CAL formation

Univariate, linear mixed model and multivariate logistic regression (LR) analyses. In the first study, no variables were associated with CAL formation. However, the linear mixed model revealed that the mean u-β2MG value \( (p = 0.034) \) and deviation value of C-reactive protein (CRP; \( p = 0.002 \)) were positively associated with CAL formation. Furthermore, the mean aspartate aminotransferase (AST) level positively associated with CALs \( (p = 0.064) \). The second study showed that the maximum pre-treatment u-β2MG value \( (p = 0.0175) \) and the occurrence of relapse \( (p = 0.0175) \) were significantly associated with CALs and the maximum pre-treatment CRP value was positively associated with CALs \( (p = 0.0855) \). The maximum pre-treatment u-β2MG and CRP values were selected as potential risk factors associated with CAL formation on the basis of the common characteristics of the first and second studies. The LR analysis demonstrated that maximum u-β2MG and maximum coronary diameter before treatment were positively and significantly associated factors \( (p = 0.063 < 0.0001) \).

Mean structure equation model for path models. Path models were built using age and sex as clinical backgrounds, IVIGR and number of IVIG treatment courses as factors associated with KD, and the maximum pre-treatment u-β2MG and CRP values as elucidated factors.

The path model in the first and second studies showed an excellent fit based on the following values: root mean square error of approximation (RMSEA) of \(< 0.0001\) and \(< 0.0001\); comparative fit index (CFI) of \(1.000 < 1.000\); and standardised path coefficient of the latent variable in terms of total effects on CALs as 0.80 and 0.76 in the first and second studies, respectively (Fig. 2a,b).

Discrimination of CALs using the SS. In the first and second studies, the SSs for the latent variable were significantly higher in patients with CALs than in those without them \( (p = 0.002; 0.063 < 0.0001) \). The receiver-operating characteristic (ROC) analyses revealed that the cutoff SSs to discriminate the development of CALs were 2.0 and 2.1 in the first and second studies, respectively.

To confirm whether the potential value of SS for discriminating CALs was true, we integrated all the data in the third study. In addition, an SEM analysis revealed a good fit based on the following values: RMSEA of \(0.032; 0.0001\); CFI of \(0.985 < 0.0001\); and standardised path coefficient of the latent variable in terms of total effects on CALs as \(0.80 < 0.76\) in the first and second studies, respectively (Fig. 2c).

The dotted and bold paths denote the differences between (a,c) and between (b,c), respectively. CAL coronary artery lesion, CRP C-reactive protein, u-β2MG urinary β2-microglobulin, IVIGR intravenous immunoglobulin resistance, IVMP times number of intravenous methylprednisolone pulse treatment courses, e error.
Figure 3. Comparison of sample scores between the patients with CALs and those without CALs in each study. (a) First study. (b) Second study. (c) Third study. The statistical analysis was performed using the Student’s t test. CAL, coronary artery lesion.

Figure 4. Comparison of sample scores between the patients with CALs and those without CALs when CAL was defined as 3.0 SD (a), 2.5 SD (b) and 2.0 SD (c). The sample scores for the latent variable discriminated CALs best when CAL was defined as 3.0 SD. The numbers in parentheses indicate the numbers of patients. Student’s t test was used for the statistical analysis. CAL coronary artery lesion.

CALs based on different definitions reduced in a dose-dependent manner according to the criteria of CALs from 3.0 to 2.0 SD (Fig. 4).

To investigate the association of the latent factor with the maximum coronary artery diameters upon admission, another SEM analysis including the maximum values of the coronary artery diameters upon admission was performed. The analysis revealed no direct association between the latent variable and maximal diameters of the coronary arteries upon admission (standardised path coefficient, 0.095; \( p = 0.25 \)).
The Nnet analysis included seven input variables comprising the maximum diameters of the coronary arteries on admission and six other variables used in the SEM analysis. All these variables were normalized as previously described. The output variable was the SS for the development of CALs (Fig. 5). Subsequently, we obtained a statistical model with four nodes in the intermediate layer, an overfit penalty of 0.02, $R^2$ of 0.89 and $R^2$ of 0.64 by the five-fold cross-validation method, although there were a few outlier patient (SI Fig. S1). The sensitivity and specificity for the prediction of CAL development were 73% and 99%, respectively, when the cutoff SS value was 2.0.

The combined data of LR and Nnet analyses did not increase the sensitivity and specificity of 73% and 99%, respectively (Table 2). In a new data set consisting of 38 KD patients from 2019 to 2020 at our hospital, the sensitivity and specificity were 50% and 100%, respectively.

Discussion. We developed a new method of predicting CALs in KD using SEM and Nnet analyses. This is the first study to show the causal associations between a latent factor of CALs and the presence of CALs. Several articles have dealt with the prediction of CALs; some of which presented laboratory data on special substances like NT-proBNP or a cytokine of HGF, whereas others focused on clinical data. The method used in this study is superior to all the previous methods described in former studies. This is because it is the first to indicate the usefulness of SS, a latent variable, predicting CALs in KD patients. Although the predicting ability in this case was slightly inferior to that of the previous report in terms of c-index, we believe that there are some advantages compared to the previous reports. First, the predicting ability of LR analyses using measurable variables was fair in previous studies, whereas it was poor in this case. We could attribute this to racial differences. Similar discrepancies were often highlighted in the risk score for IVIGR in KD patients. In our study, the SS used in Nnet analysis was a latent factor obtained by AMOS. Moreover, the latent factor was supposed to be a reflection of the predisposition to KD patients; thus, some genes associated with the predisposition might be more important than the real measured data in the Japanese population for the prediction of CALs in KD. Further studies are necessary to elucidate the details.

Second, measured only three parameters in daily clinical practice, i.e. serum CRP level, u-β2MG/Cr ratio, and the maximal diameters of the coronary arteries upon admission. The other necessary information for our method was easily obtained upon admission like age, sex, IVIGR (Sano score) and number of IVMP treatment courses due to the risk based on IVIGR scores.

Third, we agree that a sensitivity of 73% and specificity of 99% with a c-index of 0.86 is not exceptional; however, this is thought to be better than the IVIGR scores, because we can directly predict the formation of CALs instead of formulating the clinical process of IVIGR patients. Since we found some outlier patients in the analysis, it is speculated that the sensitivity can be increased if we find more appropriate factors for the input layers of Nnet analysis. Further studies are needed in the future.

This study included data from two retrospective studies. In the first study, no significant differences between the investigated parameters were observed. This was supposed to be derived from the different measurers of

**Table 2.** Comparison of sensitivity, specificity and c-index among the three statistical models. The numbers in parentheses indicate the number of patients.

|                          | (A) Logistic regression analysis model data | (B) Neural networks analysis model data | Combined A and B model data |
|--------------------------|-------------------------------------------|----------------------------------------|----------------------------|
| Sensitivity              | 36% (4/11)                                | 73% (8/11)                             | 73% (8/11)                 |
| Specificity              | 99% (232/234)                             | 99% (232/234)                          | 99% (231/234)              |
| C-index                  | 0.68                                      | 0.86                                   | 0.86                       |

**Neural network analysis.** The Nnet analysis included seven input variables comprising the maximum diameters of the coronary arteries on admission and six other variables used in the SEM analysis. All these variables were normalized as previously described. The output variable was the SS for the development of CALs (Fig. 5). Subsequently, we obtained a statistical model with four nodes in the intermediate layer, an overfit penalty of 0.02, $R^2$ of 0.89 and $R^2$ of 0.64 by the five-fold cross-validation method, although there were a few outlier patient (SI Fig. S1). The sensitivity and specificity for the prediction of CAL development were 73% and 99%, respectively, when the cutoff SS value was 2.0.

The combined data of LR and Nnet analyses did not increase the sensitivity and specificity of 73% and 99%, respectively (Table 2). In a new data set consisting of 38 KD patients from 2019 to 2020 at our hospital, the sensitivity and specificity were 50% and 100%, respectively.

**Discussion.** We developed a new method of predicting CALs in KD using SEM and Nnet analyses. This is the first study to show the causal associations between a latent factor of CALs and the presence of CALs.

Several articles have dealt with the prediction of CALs; some of which presented laboratory data on special substances like NT-proBNP or a cytokine of HGF, whereas others focused on clinical data. The method used in this study is superior to all the previous methods described in former studies. This is because it is the first to indicate the usefulness of SS, a latent variable, predicting CALs in KD patients. Although the predicting ability in this case was slightly inferior to that of the previous report in terms of c-index, we believe that there are some advantages compared to the previous reports. First, the predicting ability of LR analyses using measurable variables was fair in previous studies, whereas it was poor in this case. We could attribute this to racial differences. Similar discrepancies were often highlighted in the risk score for IVIGR in KD patients. In our study, the SS used in Nnet analysis was a latent factor obtained by AMOS. Moreover, the latent factor was supposed to be a reflection of the predisposition to KD patients; thus, some genes associated with the predisposition might be more important than the real measured data in the Japanese population for the prediction of CALs in KD. Further studies are necessary to elucidate the details.

Second, measured only three parameters in daily clinical practice, i.e. serum CRP level, u-β2MG/Cr ratio, and the maximal diameters of the coronary arteries upon admission. The other necessary information for our method was easily obtained upon admission like age, sex, IVIGR (Sano score) and number of IVMP treatment courses due to the risk based on IVIGR scores.

Third, we agree that a sensitivity of 73% and specificity of 99% with a c-index of 0.86 is not exceptional; however, this is thought to be better than the IVIGR scores, because we can directly predict the formation of CALs instead of formulating the clinical process of IVIGR patients. Since we found some outlier patients in the analysis, it is speculated that the sensitivity can be increased if we find more appropriate factors for the input layers of Nnet analysis. Further studies are needed in the future.

This study included data from two retrospective studies. In the first study, no significant differences between the investigated parameters were observed. This was supposed to be derived from the different measurers of

**Figure 5.** Artificial neural networks for predicting CALs using sample scores. Intermediate variable (H1) was denoted as the formula shown in SI Table S1. IVIGR intravenous immunoglobulin resistance, IVMP times number of intravenous methylprednisolone pulse treatment courses, CRP C-reactive protein, u-β2MG urinary β2-microglobulin, SS sample score.
coronary artery diameters among the patients. However, the use of a mixed linear model eliminated this problem and revealed that the maximal serum CRP level and u-β2MG/Cr ratio were statistically significant factors because the analysis reduced the degree of repeated measurement errors. In the second study, the same measurer obtained the coronary artery diameters that enabled us to identify the important parameters using the Student’s t test and chi-square test. These results confirmed the importance of standardisation of coronary artery measurements. Several reports have indicated that maximal CRP values are associated with CAL formation\(^1\). In contrast, no report has identified the importance of u-β2MG/Cr values. However, these findings might be supported by the observation that suggests the importance of u-β2MG/Cr values for IVIGR\(^2\) because IVIGR is a risk factor of CAL formation\(^3\). β2MG is involved in the cytokine networks\(^4\); therefore, the relationship between u-β2MG/Cr values and the development of KD needs to be investigated, particularly from the point of view of abnormal innate immunity, which is associated with KD\(^7\).

Recently, several genes have been associated with KD\(^5\), suggesting the presence of unknown factors associated with its pathogenesis. Using standard statistical methods, the effects of unknown factors could be investigated. Thus, we used the SEM analysis to clarify the role of a latent factor in CAL formation.

The SEM was devised in the 1970s\(^3\) and was routinely used for analysing social and psychological aspects of problems\(^7\) because it involves several complicated and unknown factors. However, we sometimes encounter the complicated relationships between the clinical parameters even in the field of paediatric clinical medicine, suggesting that the SEM model may be suitable in the analysis of statistical data obtained from our daily clinical practice.

In fact, using SEM analysis, we clarified the role of serum IgG3 levels in the clinical severity of pandemic influenza H1N1pdm, an unknown latent factor\(^4\). Therefore, we used the SS as a latent variable in this study because it reflects the influence of the pathogenesis of KD on each patient when we set a path from the latent variable to CAL formation. Since the efficiency of CAL discrimination by SS was associated with the cutoff SS by the ROC analysis, we thought that the SS can be used for predicting CAL formations in KD patients. Indeed, our hypothesis was proven true by the fact that CAL formation was remarkably discriminated by the SS (\(p < 0.0001\)). In addition, the presence of the causal association between the SS and CAL formation was confirmed by our finding that the potential value of SS to discriminate CALs based on the different definitions reduced in a dose-dependent manner according to the criteria for CAL from 3.0 to 2.0 SD.

Based on our data, we speculate that unknown factors possibly relevant to predisposition, representative of latent variables, are important for CAL formation in Japanese KD patients. To prove this hypothesis, we eliminated a latent factor that led to a severe reduction in the \(R^2\) value for CALs from 1.000 to 0.020 (data not shown). This also suggests the importance of the SEM analysis in creating a new statistical model for predicting CAL formation at least in Japan.

In addition, no evidence of association was found in the additional SEM analysis between a latent variable and the maximal diameters of the coronary arteries upon admission, suggesting the possibility of another mechanism being involved in the previously reported association of the maximal coronary artery diameters upon admission with CAL formation\(^8\). Further studies are required on this issue.

Recently, the Nnet analysis was reported to be useful in predicting several events\(^3\). In fact, we previously reported on this aspect of Nnet, using it to predict future bone mineral density and annual bone loss rate in postmenopausal women\(^9\). This experience led us to use Nnet analysis to predict the occurrence of CALs in the third study. The analysis enabled us to develop a good and simple statistical model for \(R^2 = 0.89\) and \(R^2 = 0.64\) in the five-fold cross-validation study to predict CALs with a sensitivity and specificity of 73% and 99%, respectively, using a cutoff SS of 2.0. The five-fold cross-validation study is proposed to reduce the possibility of overfitting, which is recommended in a clinical study using relatively small number of patients. However, this method is not perfect.

To confirm whether our methods have translational impacts, we investigated a new data set involving 38 KD patients from 2019 to 2020 at our hospital, including two patients transferred to the tertiary referral hospital because we had no KD patients with CALs from 2019 to 2020. Using the combined method of Nnet and LR analyses, it revealed sensitivity and specificity of 50% and 100%, respectively. These data did not contradict the retrospective model data in this study, despite the few patients. On the basis of these data, we believe that our protocols for predicting CALs in KD patients using the Nnet analysis have translational impacts on the field of clinical paediatrics.

For the Nnet analysis, we added the maximal coronary artery diameters upon admission to the input variables. This is supported by a previous report\(^10\) that suggested the important role of the maximal coronary artery diameters upon admission for the accurate prediction of CAL formation.

The influence of this study’s IVMP design on the evaluation of sensitivity and specificity may be questioned. To answer this, we investigated the effect of elimination of IVMP in the AMOS study, which increased the value of SS to 2.3. Subsequently, we performed Nnet analysis again to select the new program and found cross-validation values of \(R^2 = 0.84\) and 0.66. It demonstrated a sensitivity and specificity of 55% and 99%, respectively, and the former value was < 73% in this study. These data suggested the influence of IVMP therapy on the prediction of CALs. We speculated the relationship between these findings and the evidence of the usefulness of IVMP to prevent CALs.

The timing of the factors to sample and input is another aspect to consider. For u-β2MG/Cr and CRP values, we used the maximal values before starting the first-line therapy in our hospital. For the diameter of coronary arteries, we used the values before starting the first-line therapy and also the maximal values during admission in our hospital. For the new data set, we added the two patients in the second study with CALs based on the reports from the tertiary referral hospital. For the needed number of IVMP treatments, we are able to decide based on IVIGR scores on admission. Therefore, we can decide all the input variables for this method before the first-line therapy. Thus, our method can help the doctors predict the occurrence of CALs upon admission and decide whether they should opt for the second- or third-line therapy for KD earlier. Also, we conclude that
the findings in this study are necessary in the clinical setting as the results predicted CALs to have a sensitivity, specificity and c-index of 73%, 99% and 0.86, respectively.

There are several limitations of using the Nnet method. One is the problem of overfitting. To avoid this, we used a five-fold cross-validation method to select a program when $R^2$ values on cross-validation were $> 0.64$ ($r = 0.8$). Moreover, we investigated a new data set to reconfirm the translational impacts of our method on the field of paediatric clinics. However, these data need to be further investigated because the new data set had fewer patients. Another limitation is the possible presence of other suitable variables for the input layer of Nnet because of the presence of few outlier patients to predict SS for CALs.

There are also some limitations to using the AMOS method. One is the SS that was used to predict CALs. Since the SS was calculated from the data of each sample with the mean value and deviations from the mean by AMOS, SS might differ with hospitals, depending on the clinical severity of KD patients and the clinical role of each hospital, such as that of tertiary hospitals.

Another limitation could be the treatment method, because its process is different for patients from March 2002 to 2005 and patients from July 2008 to April 2012 studies, even though we tried to correct the bias statistically through AMOS and Nnet. This limitation is thought to be derived using a retrospective study design and could be clarified by a prospective study in the future.

Finally, a future confirmatory study with a large number of Japanese patients using these methods is necessary since the incidence of CALs in this study was approximately 5%, which resulted in only 14 KD patients and CALs for analyses.

In conclusion, we developed a good and simple statistical model with remarkable accuracy in discriminating CALs using common parameters that are used in the field of clinical medicine. This method is useful for deciding the time and type of therapy for Japanese KD patients to prevent CAL formation in the future.

**Methods**

We conducted a retrospective investigation using the opt-out method of our hospital. The ethics review board of the Minoh City Hospital approved the study and also waived the need for informed consent. All methods were performed per the Declaration of Helsinki and relevant guidelines.

**Subjects**

We included 375 children (202 boys, 152 girls and 21 others (gender not known)) who were clinically diagnosed with KD and admitted to the Minoh City Hospital between March 2002 and 2005, between July 2008 and April 2012 (the first study) and between July 2014 and December 2018 (the second study). Patients with $< 4$ of the diagnostic criteria for KD ($n = 29$), patients who were transferred to a tertiary referral hospital ($n = 16$), or had serious complications ($n = 16$) like anaphylaxis and drug-induced hypersensitivity syndrome were excluded. Finally, 314 children (185 boys and 129 girls) were deemed eligible for this study. Patients with at least two of the following three variables were classified into the high-risk group who might be non-responsive to IVIG therapy: CRP level $\geq 7$ mg/dL, total bilirubin (T-Bill) level $\geq 0.9$ mg/dL and aspartate aminotransferase (AST) level $\geq 200$ U/L (Sano score)$^{14}$. The remaining patients were classified into the low-risk group. In the first study, we used IVIG (2 g/kg) with IVMP (30 mg/kg) therapy for patients in the high-risk group ($n = 13$) and only IVIG therapy for those in the low-risk group ($n = 34$) between March 2002 and 2005. On the other hand, we used IVIG with two doses of IVMP therapy for patients in the high-risk group ($n = 17$) and IVIG with IVMP for those in the low-risk group ($n = 42$) between July 2008 and April 2012. In the second study, some patients with CRP levels of $\geq 7$ mg/dL were treated with IVIG with IVMP therapy ($n = 113$) and those with CRP levels of $< 7$ mg/dL were treated with only IVIG therapy in the low-risk group ($n = 75$), whereas those in the high-risk group ($n = 20$) were treated with IVIG with two doses of IVMP therapy (Fig. 1).

**Echocardiography and laboratory tests.** To evaluate coronary artery dilation, two-dimensional echocardiographic measurements of coronary artery diameters were performed. In detail, the diameters of the left main coronary trunk artery (LMT), proximal left anterior descending coronary artery (LAD) and proximal right coronary artery (RCA) were measured at three time points: before the treatment, immediately after the treatment and at hospital discharge. The diameters of the coronary arteries were expressed as $z$ scores calculated using the least mean square method.$^{36}$ We defined CALs as a maximum $z$ score of $\geq 3.0$ for the LMT, LAD or RCA. Laboratory tests were performed at the above mentioned three time points. Serum constituents were measured with the standard techniques used at Minoh City Hospital. The levels of u-$\beta$2MG were measured using a latex immunoassay kit (Wako, Osaka, Japan).

**Study design.** We performed a retrospective evaluation of KD patients to elucidate the risk factors of CAL formation. The investigated variables were age; sex; white blood cell count; levels of CRP, T-Bill, AST and u-$\beta$2MG/Cr; presence of relapse; IVIGR (Sano score) and number of IVMP treatment courses. In the first study, the diameters of the coronary arteries were measured by several physicians, which may have caused a measurement variation in addition to several missing clinical data. Therefore, we used a linear mixed model in the first study. In contrast, we used the Student’s $t$ test and chi-square test in the second study because the coronary artery diameters were measured by only one physician with few missing clinical data. We selected variables to identify the risk factors of CAL formation from the first and second studies in terms of the statistical significance and tendency of the common characteristics of CALs. On the other hand, we investigated the possibility of usefulness of neutrophil counts, sodium and albumin by logistic regression analyses in our preliminary studies but we did not find significant effects on predicting CALs. In this investigation, we did not include hematocrit because we were not able to find it as the predictor of the occurrence of CALs by the tool of pubmed central. The SEM was used to
build path models with a latent variable. In the first study, paths between the variables were selected when they showed a significant correlation of \( p < 0.002 \) with correlation coefficients of \( > 0.3 \). In the second study, we selected the path that showed the coefficient values of the maximal standardised path between a latent variable and CALs on the basis of a significant correlation of \( p < 0.0001 \), with correlation coefficients of \( > 0.3 \). Finally, we integrated all the data of the first and second studies to investigate the possibility of predicting the development of CALs using the SEM (the third study). In the third study, the criteria for significant correlation of \( p < 0.0001 \) with correlation coefficients \( > 0.33 \) were used. We supposed that the correlation coefficient no less than 0.3 is clinically more important than the \( p \) values. This explains why the cutoff \( p \) values were different among the first and second studies. In all three studies, we examined the statistical significance between the development of CALs and the SSs for the latent variable using Student's \( t \) tests. Thereafter, a ROC analysis was conducted to determine the cutoff SS for discriminating the CALs.

In the Nnet, we used the data on the diameters of the coronary arteries upon admission in addition to the same variables selected as those in the SEM in the third study. Two to four nodes in the intermediate layer from 0.01, 0.02 and 0.04 overfit penalties were selected. For the selected statistical model, the criteria of the best measured \( R^2 \geq 0.81 \) (\( r = 0.9 \)) and \( R^2 \geq 0.64 \) (\( r = 0.8 \)) for five-fold cross-validation method were used.

Furthermore, LR analysis was performed using age and sex as clinical backgrounds, IVIG and number of IVIG treatment courses as factors associated with KD\textsuperscript{m} and the maximum u-β2MG, CRP values and maximal coronary diameter before the treatment as elucidated factors in the third-study patients. Furthermore, we investigated the usefulness of combined data of LR (real measured data) and NNet analyses (predicted scores by AMOS). We considered it positive when one of the analyses showed the positive occurrence of CALs.

Lastly, we investigated a new data set to confirm whether our methods have translational impacts. This set included 38 KD patients from 2019 to 2020 at our hospital. Because there were no patients with CALs in the new samples, we included two patients who were transferred to the tertiary hospital and had CALs but were not analysed in the previous study based on the exclusion criteria. One patient was treated with IVIG with two new samples, we included two patients who were transferred to the tertiary referral hospital and he was treated with plasmapheresis. The other one was treated with IVIG with IVMP therapy in Minoh City Hospital; however, fever persisted after the therapy. Then, we transferred the patient to the tertiary referral hospital and he was treated with plasmapheresis. The other one was treated with IVIG therapy in Minoh City Hospital; however, five KD symptoms persisted in spite of the therapy. Thus, we transferred the patient to a tertiary referral hospital and she was treated with additional IVIG with oral cyclosporin.

**Statistical analyses.** \( p < 0.05 \) and \( 0.05 < p < 0.1 \) were accepted as statistically significant and prone, respectively. The statistical significance of the path models was judged from the following criteria: RMSEA \( < 0.05 \), AIC score \( < 70 \), CFI score \( > 0.95 \) and \( R^2 \) score \( > 0.95 \). The JMP version 8.0 software (SAS Institute, Cary, NC) was used for the Student's \( t \) test, chi-square test and NNet analysis. AMOS 23.0 (IBM-SPSS, USA) and SPSS version 23.0 (IBM Corp, Armonk, NY, USA) were used for the SEM and linear mixed model, respectively.

**Data availability**
The datasets analysed during the current study are available from the corresponding author on reasonable request.

Received: 11 December 2019; Accepted: 28 June 2020
Published online: 17 July 2020

**References**
1. Kawasaki, T. Kawasaki disease. *Proc. Ipn. Acad. Ser. B Phys. Biol. Sci.* **82**, 59–71. https://doi.org/10.1283/pia082.59 (2006).
2. Takahashi, K., Oharaseki, T., Yokouchi, Y., Hiruta, N. & Naoe, S. Kawasaki disease as a systemic vasculitis in childhood. *Ann. Vasc. Dis.* 3, 173–181. https://doi.org/10.3400/avd.asavp01003 (2010).
3. Kikut, H. et al. Kawasaki disease and an unusual primary infection with Epstein–Barr virus. *Pediatrics* 73, 413–414 (1984).
4. Matsuhara, K. & Fukaya, T. The role of superantigens of group A Streptococcus and *Staphylococcus aureus* in Kawasaki disease. *Crit. Opin. Infect. Dis.* 20, 298–303. https://doi.org/10.1097/QCO.0b013e3280964dc8 (2007).
5. Takeshita, S. et al. The role of bacterial lipopolysaccharide-bound neutrophils in the pathogenesis of Kawasaki disease. *J. Infect. Dis.* 179, 508–512. https://doi.org/10.1086/314400 (1999).
6. Manlihot, C. et al. Environmental epidemiology of Kawasaki disease: Linking disease etiology, pathogenesis and global distribution. *PloS One* 13, e0191087. https://doi.org/10.1371/journal.pone.0191087 (2018).
7. Hara, T. et al. Kawasaki disease: A matter of innate immunity. *Clin. Exp. Immunol.* 186, 134–143. https://doi.org/10.1111/celi.12832 (2016).
8. Onouchi, Y. et al. ITPKC functional polymorphism associated with Kawasaki disease susceptibility and formation of coronary artery aneurysms. *Nat. Genet.* 40, 35–42. https://doi.org/10.1038/ng.2007.59 (2008).
9. Onouchi, Y. The genetics of Kawasaki disease. *Int. J. Rheum. Dis.* 21, 26–30. https://doi.org/10.1111/1756-185x.13218 (2018).
10. Kimura, Y. et al. Identification of candidate diagnostic serum biomarkers for Kawasaki disease using proteomic analysis. *Sci. Rep.* 7, 43732. https://doi.org/10.1038/srep43732 (2017).
11. Lo, M. S. & Newburger, J. W. Role of intravenous immunoglobulin in the treatment of Kawasaki disease. *Int. J. Rheum. Dis.* 21, 64–69. https://doi.org/10.1111/1756-185x.13220 (2018).
12. Kobayashi, T. et al. Prediction of intravenous immunoglobulin unresponsiveness in patients with Kawasaki disease. *Circulation* 113, 2606–2612. https://doi.org/10.1161/circulationaha.105.592865 (2006).
13. Egami, K. et al. Prediction of resistance to intravenous immunoglobulin treatment in patients with Kawasaki disease. *J. Pediatr.* 149, 237–240. https://doi.org/10.1016/j.jpeds.2006.03.050 (2006).
14. Sano, T. et al. Prediction of non-responsiveness to standard high-dose gamma-globulin therapy in patients with acute Kawasaki disease before starting initial treatment. *Eur. J. Pediatr.* 166, 131–137. https://doi.org/10.1007/s00431-006-0223-z (2007).
15. Rigante, D. et al. Critical overview of the risk scoring systems to predict non-responsiveness to intravenous immunoglobulin in Kawasaki Syndrome. *Int. J. Mol. Sci.* 17, 278. https://doi.org/10.3390/ijms17030278 (2016).
16. Son, M. B. F. et al. Predicting coronary artery aneurysms in Kawasaki Disease at a North American center: An assessment of baseline z scores. J. Am. Heart Assoc. https://doi.org/10.1161/jaha.116.005378 (2017).
17. Kaneko, K. et al. Prediction of the risk of coronary artery lesions in Kawasaki disease by brain natriuretic peptide. Pediatr. Cardiol. 32, 1106–1109. https://doi.org/10.1007/s00246-011-9986-8 (2011).
18. Okuma, Y. et al. Serum tenascin-C as a novel predictor for risk of coronary artery lesion and resistance to intravenous immunoglobulin in Kawasaki disease—a multicenter retrospective study. Circ. J. 80, 2376–2381. https://doi.org/10.1253/circj.CJ-16-0565 (2016).
19. Ohno, T. et al. Serum hepatocyte growth factor combined with vascular endothelial growth factor as a predictive indicator for the occurrence of coronary artery lesions in Kawasaki disease. Eur. J. Pediatr. 161, 105–111. https://doi.org/10.1007/s00431-001-0860-1 (2002).
20. Bai, L. et al. Retrospective analysis of risk factors associated with Kawasaki disease in China. Oncotarget 8, 54357–54363. https://doi.org/10.18632/oncotarget.17530 (2017).
21. Son, M. B. F. et al. Risk model development and validation for prediction of coronary artery aneurysms in Kawasaki disease in a North American population. J. Am. Heart Assoc. 8, e011319. https://doi.org/10.1161/jaha.118.011319 (2019).
22. Kobayashi, T. et al. Efficacy of immunoglobulin plus prednisolone for prevention of coronary artery abnormalities in severe Kawasaki disease (RAISE study): A randomised, open-label, blinded-endpoints trial. Lancet 379, 1613–1620. https://doi.org/10.1016/s0140-6736(11)61930-2 (2012).
23. Miyata, K. et al. Efficacy and safety of intravenous immunoglobulin plus prednisolone therapy in patients with Kawasaki disease (Post RAISE): A multicentre, prospective cohort study. Lancet Child Adolesc. Health 2, 855–862. https://doi.org/10.1016/s2352-4642(18)30293-1 (2018).
24. Okada, K. et al. Pulse methylprednisolone with gammaglobulin as an initial treatment for acute Kawasaki disease. Eur. J. Pediatr. 168, 181–185. https://doi.org/10.1007/s00431-008-0727-9 (2009).
25. Okubo, Y. et al. Association between dose of glucocorticoids and coronary artery lesions in Kawasaki disease. Arthritis Care Res. 70, 1052–1057. https://doi.org/10.1002acr.23456 (2018).
26. Adya, M. & Collopy, F. How effective are neural networks at forecasting and prediction? A review and evaluation. J. Forecast. 17, 481–495. https://doi.org/10.1002/(issn)1099-131x(199809017)5:6<481::aid-for709>3.0.co;2-q (1998).
27. Jone, P. N. et al. Infliximab plus intravenous immunoglobulin (IVIG) versus IVIG alone as initial therapy in children with Kawasaki disease presenting with coronary artery lesions: Is dual therapy more effective? Pediatr. Infect. Dis. J. 37, 976–980. https://doi.org/10.1097/infd.0000000000001951 (2018).
28. Hamada, H. et al. Efficacy of primary treatment with immunoglobulin plus ciclosporin for prevention of coronary artery abnormalities in patients with Kawasaki disease predicted to be at increased risk of non-response to intravenous immunoglobulin (KAICA): A randomised controlled, open-label, blinded-endpoints, phase 3 trial. Lancet 393, 1128–1137. https://doi.org/10.1016/s0140-6736(18)32003-8 (2019).
29. Mori, M. et al. Efficacy of plasma exchange therapy for Kawasaki disease intractable to intravenous gamma-globulin. Mod. Rheumatol. 14, 43–47. https://doi.org/10.1007/s10165-003-0264-3 (2004).
30. Shioji, M. et al. Artificial neural networks to predict future bone mineral density and bone loss rate in Japanese postmenopausal women. BMC Res. Notes 10, 590. https://doi.org/10.1186/s13104-017-2910-4 (2017).
31. Yoshikawa, T. et al. Urinary β2 microglobulin to creatinine ratio as a predictive factor of intravenous immunoglobulin resistance. Circulation 131 suppl 2, A107. https://doi.org/10.1161/circulationaha.1131.suppl_116.1107 (2015).
32. Li, L., Dong, M. & Wang, X. G. The implication and significance of beta 2 microglobulin: A conservative multifunctional regulator. Chin. Med. J. 129, 448–455. https://doi.org/10.4103/0366-6999.176084 (2016).
33. Tarka, P. An overview of structural equation modeling: Its beginnings, historical development, usefulness and controversies in the social sciences. Qual. Quant. 52, 313–354. https://doi.org/10.1007/s11135-017-0469-4 (2018).
34. Yamamoto, T. et al. Serum immunoglobulin G subclass levels and estimated clinical severity caused by possible influenza A (H1N1) pdm 2009 infection. J. Infect. Chemother. 19, 833–842. https://doi.org/10.1016/j.jiac.2013.05.015 (2013).
35. Lisboa, P. J. A review of evidence of health benefit from artificial neural networks in medical intervention. Neural Netw. 15, 11–39 (2002).
36. Cole, T. J. & Green, P. J. Smoothing reference centile curves: The LMS method and penalized likelihood. Stat. Med. 11, 1305–1319. https://doi.org/10.1002/sim.478011005 (1992).

Acknowledgements
The authors thank Masashi Mukai, Yuri Kasui, Takafumi Iguchi, Takeshi Ishimi, Ryuko Kora, Toshiya Katsura and Hidehiro Suginobe who were attending physicians of KD patients admitted to Minoh City Hospital and contributed to data collection in this study. We also thank Tadamori Takahara who was attending physicians of KD patients transferred from our hospital to Hyogo Prefectural Amagasaki General Medical Center.

Author contributions
T.Y. conceptualised and designed the study and performed the statistical analysis for SEM and Nnet. J.A. contributed to the conceptualisation and design of the study. T.Y. and J.A. wrote the manuscript. M.N., Y.H., E.K., T.S. and Y.M. contributed to the data collection and interpretation. All authors approved the submission of the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-68657-0.

Correspondence and requests for materials should be addressed to T.Y.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
