Hybrid methods for elliptic and hyperbolic PDEs

<u>Victor Michel-Dansac</u>^{*}, joint work with Hélène Barucq[†], Michel Duprez[‡], Florian Faucher[†], Emmanuel Franck^{*}, Frédérique Lecourtier[‡], Vanessa Lleras[§], Laurent Navoret^{*}, Nicolas Victorion[†]

Monday, December 02, 2024 Workshop MEDIA, La Garde

*<u>MACARON project-team, Université de Strasbourg, CNRS, Inria, IRMA, France</u> †MAKUTU project-team, Université de Pau et des Pays de l'Adour, CNRS, Inria, France ‡MIMESIS project-team, Université de Strasbourg, CNRS, Inria, France §IMAG, Université de Montpellier, France





Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

To fix notation, consider the following stationary PDE:

$$\begin{cases} \mathcal{D}(W, x) = 0 & \text{ for } x \in \Omega, \\ W(x) = g(x) & \text{ for } x \in \partial\Omega, \end{cases}$$

where

- $\Omega \subset \mathbb{R}^d$ is the spatial domain,
- $W \in \mathbb{R}^q$ is the unknown solution,
- + $\ensuremath{\mathfrak{D}}$ is some differential operator,
- g is a known function.

Parametric approximation in some classical methods for stationary PDEs

In many classical numerical methods, the solution is approximated by a parametric function, **linear in its parameters**, and a basis $(\varphi_i)_i$ depending on the chosen method:

$$W_{\theta}(x) = \sum_{j=1}^{N} \theta_{j} \varphi_{j}(x);$$
 the parameters $(\theta_{j})_{j \in \{1,...,N\}}$ are called degrees of freedom.

Parametric approximation in some classical methods for stationary PDEs

In many classical numerical methods, the solution is approximated by a parametric function, **linear in its parameters**, and a basis $(\varphi_j)_j$ depending on the chosen method:

$$W_{\theta}(x) = \sum_{j=1}^{N} \theta_j \varphi_j(x);$$
 the parameters $(\theta_j)_{j \in \{1,...,N\}}$ are called degrees of freedom.

- <u>mesh-based</u> methods, with mesh $(x_j)_{j \in \{1,...,N\}}$:
- finite difference: $\varphi_j = \delta_{x_j}$
- **1**st-order finite volume: φ_j are piecewise constant
- **finite element**: φ_j are piecewise polynomial

Parametric approximation in some classical methods for stationary PDEs

In many classical numerical methods, the solution is approximated by a parametric function, **linear in its parameters**, and a basis $(\varphi_j)_j$ depending on the chosen method:

 $W_{\theta}(x) = \sum_{j=1}^{N} \theta_{j} \varphi_{j}(x);$ the parameters $(\theta_{j})_{j \in \{1,...,N\}}$ are called degrees of freedom.

- <u>mesh-based</u> methods, with mesh $(x_j)_{j \in \{1,...,N\}}$:
- finite difference: $\varphi_j = \delta_{x_j}$
- **1**st-order finite volume: φ_j are piecewise constant
- **Finite element**: φ_j are piecewise polynomial

- <u>mesh-free</u> methods:
- **spectral methods**: $\varphi_j = e^{ik_jx}$ in Fourier space
- ► **SPH**: $\varphi_j(x) = \Xi(|x x_j|)$ with Ξ a kernel function
- **diffuse elements**: φ_j are piecewise polynomial

Parametric approximation in some classical methods for stationary PDEs

In many classical numerical methods, the solution is approximated by a parametric function, **linear in its parameters**, and a basis $(\varphi_i)_i$ depending on the chosen method:

 $W_{\theta}(x) = \sum_{j=1}^{N} \theta_{j} \varphi_{j}(x);$ the parameters $(\theta_{j})_{j \in \{1,...,N\}}$ are called degrees of freedom.

- <u>mesh-based</u> methods, with mesh $(x_j)_{j \in \{1,...,N\}}$:
- finite difference: $\varphi_j = \delta_{x_j}$
- **1**st-order finite volume: φ_j are piecewise constant
- **finite element**: φ_j are piecewise polynomial

- <u>mesh-free</u> methods:
- **spectral methods**: $\varphi_j = e^{ik_jx}$ in Fourier space
- **SPH**: $\varphi_j(x) = \Xi(|x x_j|)$ with Ξ a kernel function
- **diffuse elements**: φ_j are piecewise polynomial

Most of these approaches are **local in space**, and the number of degrees of freedom *N* **exponentially increases with the dimension**.

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad \forall \psi \in \mathcal{H}^1_0(\Omega), \quad \int_\Omega \nabla W \cdot \nabla \psi \, dx = \int_\Omega f \psi \, dx. \end{cases}$$

<

Consider the **Poisson problem** and its **weak formulation**, with unknown $W \in \mathcal{H}^1_0(\Omega)$:

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial \Omega, \end{cases} \qquad \Longleftrightarrow \qquad \forall \psi \in \mathcal{H}^1_0(\Omega), \quad \int_\Omega \nabla W \cdot \nabla \psi \, dx = \int_\Omega f \psi \, dx. \end{cases}$$

Now, approximate $\mathcal{H}_0^1(\Omega)$ by a **linear subspace** of polynomial functions $V = \text{Span}((\varphi_j)_j)$.

The finite element approximation of W is, for $x \in \Omega$, $W_{\theta}(x) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$.

Now, approximate $\mathcal{H}_0^1(\Omega)$ by a **linear subspace** of polynomial functions $V = \text{Span}((\varphi_j)_j)$. The **finite element approximation** of W is, for $x \in \Omega$, $W_{\theta}(x) = \sum_{j=1}^N \theta_j \varphi_j(x)$. Plugging these approximations in the weak formulation, we get

$$\forall k \in \{1,\ldots,N\}, \quad \sum_{j=1}^{N} \theta_{j} \underbrace{\int_{\Omega} \nabla \varphi_{j} \cdot \nabla \varphi_{k} \, dx}_{A_{kj}} = \underbrace{\int_{\Omega} f \varphi_{k} \, dx}_{b_{k}},$$

i.e., with $\theta = (\theta_j)_j$, $A = (A_{kj})_{kj}$ and $b = (b_k)_k$, we have the **linear system** $A\theta = b$.

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}_0^+(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right].$$

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}^1_0(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right].$$

Now, approximate $\mathcal{H}_0^1(\Omega)$ by a **linear subspace** of polynomial functions $V = \text{Span}((\varphi_i)_i)$.

The **finite element approximation** of *W* is, for $x \in \Omega$, $W_{\theta}(x) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$. Therefore,

$$W_{\theta} = \operatorname*{argmin}_{\phi \in V} \left[\frac{1}{2} \int_{\Omega} |\nabla \phi|^2 - \int_{\Omega} f \phi \right].$$

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}^1_0(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right].$$

Now, approximate $\mathcal{H}_0^1(\Omega)$ by a **linear subspace** of polynomial functions $V = \text{Span}((\phi_i)_i)$.

The **finite element approximation** of *W* is, for $x \in \Omega$, $W_{\theta}(x) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$. Therefore,

$$\theta = \underset{\vartheta \in \mathbb{R}^{N}}{\operatorname{argmin}} \left[\sum_{j=1}^{N} \sum_{k=1}^{N} \frac{1}{2} \vartheta_{j} \vartheta_{k} \int_{\Omega} \nabla \varphi_{j} \cdot \nabla \varphi_{k} - \sum_{j=1}^{N} \vartheta_{j} \int_{\Omega} f \varphi_{j} \right].$$

We can write $\theta = \operatorname{argmin}_{\vartheta \in \mathbb{R}^N} \mathcal{J}(\vartheta)$, with \mathcal{J} a quadratic function.

Solving this **quadratic minimization problem**, we obtain the same **linear system** $A\theta = b$.

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}^1_0(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right].$$

Now, approximate $\mathcal{H}_0^1(\Omega)$ by a linear subspace of polynomial functions $V = \text{Span}((\varphi_j)_j)$.

The **finite element approximation** of *W* is, for $x \in \Omega$, $W_{\theta}(x) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$. Therefore,

$$\theta = \underset{\vartheta \in \mathbb{R}^{N}}{\operatorname{argmin}} \left[\sum_{j=1}^{N} \sum_{k=1}^{N} \frac{1}{2} \vartheta_{j} \vartheta_{k} \int_{\Omega} \nabla \varphi_{j} \cdot \nabla \varphi_{k} - \sum_{j=1}^{N} \vartheta_{j} \int_{\Omega} f \varphi_{j} \right].$$

We can write $\theta = \operatorname{argmin}_{\vartheta \in \mathbb{R}^N} \mathcal{J}(\vartheta)$, with \mathcal{J} a quadratic function.

Solving this **quadratic minimization problem**, we obtain the same **linear system** $A\theta = b$.

Example: the "Deep Ritz" method [W. E and B. Yu (2018)]

Consider the Poisson problem and its energy formulation:

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}_0^1(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right].$$

Example: the "Deep Ritz" method [W. E and B. Yu (2018)]

Consider the Poisson problem and its energy formulation:

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}_0^1(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right]. \end{cases}$$

We approximate $\mathcal{H}_0^1(\Omega)$ by the subspace $V = \{x \mapsto \varphi(x, \theta), \theta \in \mathbb{R}^N\}$, with $\varphi : \mathbb{R}^d \times \mathbb{R}^N \to \mathbb{R}$ a **nonlinear function** of both inputs.

The **nonlinear approximation** of *W* becomes, for $x \in \Omega$, $W_{\theta}(x) = \varphi(x, \theta)$. Therefore,

$$W_{\theta} = \underset{\varphi \in V}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \varphi(x, \theta)|^2 \, dx - \int_{\Omega} f(x) \varphi(x, \theta) \, dx \right].$$

Example: the "Deep Ritz" method [W. E and B. Yu (2018)]

Consider the Poisson problem and its energy formulation:

$$\begin{cases} -\Delta W = f & \text{in } \Omega, \\ W = 0 & \text{on } \partial\Omega, \end{cases} \qquad \Longleftrightarrow \qquad W = \underset{\psi \in \mathcal{H}_0^1(\Omega)}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \psi|^2 - \int_{\Omega} f \psi \right].$$

We approximate $\mathcal{H}_0^1(\Omega)$ by the subspace $V = \{x \mapsto \varphi(x, \theta), \theta \in \mathbb{R}^N\}$, with $\varphi : \mathbb{R}^d \times \mathbb{R}^N \to \mathbb{R}$ a **nonlinear function** of both inputs.

The **nonlinear approximation** of *W* becomes, for $x \in \Omega$, $W_{\theta}(x) = \varphi(x, \theta)$. Therefore,

$$\theta = \underset{\vartheta \in \mathbb{R}^{N}}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla \varphi(x, \vartheta)|^{2} dx - \int_{\Omega} f(x) \varphi(x, \vartheta) dx \right].$$

We can write $\theta = \operatorname{argmin}_{\vartheta \in \mathbb{R}^N} \mathcal{J}(\vartheta)$, with \mathcal{J} a nonquadratic function.

We now have to solve a nonlinear minimization problem!

We have presented two ways of approximating our unknown function W. In both cases, we define degrees of freedom $\theta \in \mathbb{R}^N$, and set $W(x) \simeq W_{\theta}(x) = \varphi(x, \theta)$, for all $x \in \Omega$.

The main difference lies in the **choice of the function** φ : it is always **nonlinear in space**, but its **behavior with respect to** θ changes the nature of the approximation problem.

We have presented two ways of approximating our unknown function W. In both cases, we define degrees of freedom $\theta \in \mathbb{R}^N$, and set $W(x) \simeq W_{\theta}(x) = \varphi(x, \theta)$, for all $x \in \Omega$.

The main difference lies in the **choice of the function** φ : it is always **nonlinear in space**, but its **behavior with respect to** θ changes the nature of the approximation problem.

- 1. φ is linear in θ , $\varphi(x, \theta) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$:
 - 1.1 *W* is projected onto a finite-dimensional linear subspace
 - 1.2 solve a convex quadratic optimization problem to determine θ

We have presented two ways of approximating our unknown function W. In both cases, we define degrees of freedom $\theta \in \mathbb{R}^N$, and set $W(x) \simeq W_{\theta}(x) = \varphi(x, \theta)$, for all $x \in \Omega$.

The main difference lies in the **choice of the function** φ : it is always **nonlinear in space**, but its **behavior with respect to** θ changes the nature of the approximation problem.

- 1. φ is linear in θ , $\varphi(x, \theta) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$:
 - 1.1 *W* is projected onto a finite-dimensional linear subspace
 - 1.2 solve a convex quadratic optimization problem to determine θ

- 2. φ is nonlinear in θ :
 - 2.1 W is projected onto a finite-dimensional "submanifold"
 - 2.2 solve a nonlinear, (usually) non-convex optimization problem to determine θ

We have presented two ways of approximating our unknown function W. In both cases, we define degrees of freedom $\theta \in \mathbb{R}^N$, and set $W(x) \simeq W_{\theta}(x) = \varphi(x, \theta)$, for all $x \in \Omega$.

The main difference lies in the **choice of the function** φ : it is always **nonlinear in space**, but its **behavior with respect to** θ changes the nature of the approximation problem.

- 1. φ is linear in θ , $\varphi(x, \theta) = \sum_{j=1}^{N} \theta_j \varphi_j(x)$:
 - 1.1 *W* is projected onto a finite-dimensional linear subspace
 - 1.2 solve a convex quadratic optimization problem to determine θ

- 2. φ is nonlinear in θ :
 - 2.1 W is projected onto a finite-dimensional "submanifold"
 - 2.2 solve a nonlinear, (usually) non-convex optimization problem to determine θ

Question: How to construct suitable nonlinear functions φ ?

Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

- Using PINNs for parametric PDEs
- Numerical examples
- Hybridizing the finite element method and PINNs
- Well-balanced DG methods using bases enriched with PINNs
 - Why do we need well-balanced methods?
 - Example of a physical model: the shallow water equations
 - Numerical method overview: Discontinuous Galerkin
 - Enhancing DG with Scientific Machine Learning
 - Validation
- Conclusion

Multilayer perceptron (MLP)



Schematic of an MLP (by A. Bélières-Frendo).

An MLP is a nonlinear parametric function $\varphi : \mathbb{R}^d \times \mathbb{R}^N \to \mathbb{R}^q.$

It results from a composition of several nonlinear layers. For instance, the first layer is:

$$z^1 = \sigma(\mathsf{A}^1 z^0 + b^1) \in \mathbb{R}^{q_1}$$
,

- $z^0 \in \mathbb{R}^{q_0}$ (with $q_0 = d$),
- $A^1 \in \mathcal{M}_{q_1,q_0}(\mathbb{R})$,
- $b^1 \in \mathbb{R}^{q_1}$,
- + $\sigma\in {\mathcal C}^0({\mathbb R},{\mathbb R})$, applied component-wise.

Multilaver perceptron (MLP)



Schematic of an MLP (by A. Bélières-Frendo).

An MLP is a nonlinear parametric function $\omega: \mathbb{R}^d \times \mathbb{R}^N \to \mathbb{R}^q.$

It results from a composition of several nonlinear layers. For instance, the first layer is:

$$z^1 = \sigma(A^1z^0 + b^1) \in \mathbb{R}^{q_1}$$
,

- $z^0 \in \mathbb{R}^{q_0}$ (with $a_0 = d$).
- $A^1 \in \mathcal{M}_{a_1,a_0}(\mathbb{R})$,
- $b^1 \in \mathbb{R}^{q_1}$.
- $\sigma \in C^0(\mathbb{R}, \mathbb{R})$, applied component-wise.

In the end, the function $\varphi : (z^0, \theta) \mapsto z^{\ell}$ reads

$$\varphi(z^0,\theta) = \sigma(\mathsf{A}^{\ell}\sigma(\mathsf{A}^{\ell-1}\ldots\sigma(\mathsf{A}^1z^0+b^1)\cdots+b^{\ell-1})+b^{\ell}).$$

The degrees of freedom are $\theta = (A^1, b^1, \dots, A^\ell, b^\ell) \in \mathbb{R}^N$, with $N = \sum_{i=1}^{\ell} q_i(q_{i-1}+1)$.

Universal approximation theorems

Arbitrary-width case [G. Cybenko, Math. Control Signals Systems (1989)]

Let $\sigma \in \mathbb{C}^0(\mathbb{R}, \mathbb{R})$ be a non-polynomial function. Then, for all $(m, n) \in \mathbb{N}^2$, $\mathcal{K} \subseteq \mathbb{R}^n$ compact set, $f \in \mathbb{C}^0(\mathcal{K}, \mathbb{R}^m)$, and $\varepsilon > 0$, there exist $k \in \mathbb{N}$, $A \in \mathcal{M}_{k,n}(\mathbb{R})$, $b \in \mathbb{R}^k$ and $C \in \mathcal{M}_{m,k}(\mathbb{R})$ such that

 $\|f(x) - C\sigma(Ax + b)\|_{L^{\infty}(K)} < \varepsilon.$

Arbitrary-depth case [P. Kidger and T. Lyons, (2020)]

Let $\sigma \in \mathbb{C}^0(\mathbb{R}, \mathbb{R})$ be a non-affine function, continuously differentiable in at least one point. Let $\mathbb{N}_{n,m,n+m+2}^{\sigma}$ denote the set of MLPs with *n* inputs, *m* outputs, whose hidden layers have n + m + 2 neurons, and with activation function σ . Then, for all $(m, n) \in \mathbb{N}^2$, $\mathcal{K} \subseteq \mathbb{R}^n$ compact set, $f \in \mathbb{C}^0(\mathcal{K}, \mathbb{R}^m)$, and $\varepsilon > 0$, there exists $W_{\theta} \in \mathbb{N}_{n,m,n+m+2}^{\sigma}$ such that

 $\|f(\mathbf{x}) - W_{\theta}(\mathbf{x})\|_{L^{\infty}(K)} < \varepsilon.$

Equipped with the expression of $W_{\theta} : \mathbb{R}^d \to \mathbb{R}^q$, with $W_{\theta} = \varphi(\cdot, \theta)$, the goal is to **determine the** *N* **parameters** θ such that W_{θ} is an approximation to the PDE solution *W*.

Equipped with the expression of $W_{\theta} : \mathbb{R}^d \to \mathbb{R}^q$, with $W_{\theta} = \varphi(\cdot, \theta)$, the goal is to **determine the** *N* **parameters** θ such that W_{θ} is an approximation to the PDE solution *W*.

This is done through nonlinear optimization¹: define a **loss function** \mathcal{J} measuring² the PDE residual, i.e.,

$$\mathcal{J}(\theta) = \int_{\Omega} \mathcal{D}(W_{\theta}, x)^2 \, dx + \int_{\partial \Omega} (W_{\theta}(x) - g(x))^2 \, dx.$$

The optimal parameters are then given by:

 $\theta_{\mathsf{opt}} = \operatorname*{argmin}_{\theta} \mathcal{J}(\theta).$

¹Usually, using the ADAM algorithm [D. Kingma and J. Ba, (2015)] for stochastic gradient descent. ²This corresponds to PINNs (Physics-Informed Neural Networks, M. Raissi et al., *J. Comput. Phys.* (2019)). Another way of determining parameters θ lies in the **Deep Ritz method**³. The solution remains approximated by a neural network $W_{\theta} : \mathbb{R}^d \to \mathbb{R}^q$.

This time, the PDE is written in energy form. In the case of the Poisson problem, this leads to the following minimization problem:

$$\theta = \underset{\vartheta \in \mathbb{R}^{N}}{\operatorname{argmin}} \left[\frac{1}{2} \int_{\Omega} |\nabla W_{\theta}(x)|^{2} dx - \int_{\Omega} f(x) W_{\theta}(x) dx \right].$$

We can write $\theta = \operatorname{argmin}_{\vartheta \in \mathbb{R}^N} \mathcal{J}(\vartheta)$, which is a nonlinear optimization problem.

³see W. E and B. Yu, Commun. Math. Stat. (2018)

The PINN W_{θ} approximates the solution W to the BVP:

$$\begin{cases} \mathcal{D}(W,x) = 0 & \text{ for } x \in \Omega, \\ W(x) = g(x) & \text{ for } x \in \partial\Omega. \end{cases} \xrightarrow{\sim} \begin{cases} \mathcal{D}(W_{\theta},x) \simeq 0 & \text{ for } x \in \Omega, \\ W_{\theta}(x) \simeq g(x) & \text{ for } x \in \partial\Omega. \end{cases}$$

To train the PINN (i.e., to determine the optimal parameters θ_{opt}), one fashions a loss function using the PDE residual:

$$\mathcal{J}_{\mathsf{PDE}}(\theta) = \int_{\Omega} \|\mathcal{D}(W_{\theta}, x)\|_{2}^{2} dx + \int_{\partial \Omega} \|W_{\theta}(x) - g(x)\|_{2}^{2} dx, \text{ and then } \theta_{\mathsf{opt}} = \underset{\theta}{\operatorname{argmin}} \mathcal{J}_{\mathsf{PDE}}(\theta).$$

 \rightsquigarrow How to compute the integrals?

Multidimensional integration

$$\mathcal{J}_{\mathsf{PDE}}(\theta) = \underbrace{\int_{\Omega} \|\mathcal{D}(W_{\theta}, x)\|_{2}^{2} dx}_{\mathcal{J}_{\Omega}(\theta)} + \underbrace{\int_{\partial \Omega} \|W_{\theta}(x) - g(x)\|_{2}^{2} dx}_{\mathcal{J}_{\mathsf{boundary}}(\theta)}$$

We have to compute two integrals:

- $\mathcal{J}_{\Omega}(\theta)$ over $\Omega \subset \mathbb{R}^d$,
- $\mathcal{J}_{\text{boundary}}(\theta)$ over $\partial \Omega \subset \mathbb{R}^{d-1}$.

The classical approach involves quadrature methods. However, they require a grid, which is a problem in high dimension or on complex domains...

→ Use the Monte-Carlo approach, a mesh-less method whose convergence is slow but independent of the dimension.

PINNs: advantages and drawbacks

Once trained, PINNs with Monte-Carlo integration are able to

- quickly provide an approximation to the PDE solution,
- in a mesh-less fashion and on complex domains,
- independently of the dimension.

Once trained, PINNs with Monte-Carlo integration are able to

- quickly provide an approximation to the PDE solution,
- in a mesh-less fashion and on complex domains,
- independently of the dimension.

However, PINNs

- have trouble **generalizing** to $x \notin \Omega$;
- are usually **not competitive with classical numerical methods for computational fluid dynamics**: to reach a given error (if possible), training takes longer than using a classical numerical method, and no convincing convergence results exist at the moment.

Once trained, PINNs with Monte-Carlo integration are able to

- quickly provide an approximation to the PDE solution,
- in a mesh-less fashion and on complex domains,
- independently of the dimension.

However, PINNs

- have trouble **generalizing** to $x \notin \Omega$;
- are usually **not competitive with classical numerical methods for computational fluid dynamics**: to reach a given error (if possible), training takes longer than using a classical numerical method, and no convincing convergence results exist at the moment.

The most interesting use of PINNs, in our case, is to deal with **parametric PDEs**, where dimension-insensitivity is paramount.

Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

Parametric PINNs: approximation using the PDE residual

The parametric PINN $W_{\theta}(x; \mu)$ with parameters $\mu \in \mathbb{M} \subset \mathbb{R}^{m}$ approximates the solution W to the parametric BVP:

$$\begin{cases} \mathcal{D}(W, x; \mu) = 0 & \text{for } x \in \Omega, \mu \in \mathbb{M}, \\ W(x) = g(x; \mu) & \text{for } x \in \partial\Omega, \mu \in \mathbb{M}. \end{cases} \xrightarrow{\sim} \begin{cases} \mathcal{D}(W_{\theta}, x; \mu) \simeq 0 & \text{for } x \in \Omega, \mu \in \mathbb{M}, \\ W_{\theta}(x; \mu) \simeq g(x; \mu) & \text{for } x \in \partial\Omega, \mu \in \mathbb{M}. \end{cases}$$

Parametric PINNs: approximation using the PDE residual

The parametric PINN $W_{\theta}(x; \mu)$ with parameters $\mu \in \mathbb{M} \subset \mathbb{R}^{m}$ approximates the solution W to the parametric BVP:

 $\begin{cases} \mathcal{D}(W, x; \mu) = 0 & \text{for } x \in \Omega, \mu \in \mathbb{M}, \\ W(x) = g(x; \mu) & \text{for } x \in \partial\Omega, \mu \in \mathbb{M}. \end{cases} \xrightarrow{\sim} \begin{cases} \mathcal{D}(W_{\theta}, x; \mu) \simeq 0 & \text{for } x \in \Omega, \mu \in \mathbb{M}, \\ W_{\theta}(x; \mu) \simeq g(x; \mu) & \text{for } x \in \partial\Omega, \mu \in \mathbb{M}. \end{cases}$

The loss function then becomes

$$\mathcal{J}_{\mathsf{PDE}}(\theta) = \underbrace{\int_{\mathbb{M}} \int_{\Omega} \|\mathcal{D}(W_{\theta}, x; \mu)\|_{2}^{2} dx d\mu}_{\mathcal{J}_{\Omega}(\theta)} + \underbrace{\int_{\mathbb{M}} \int_{\partial \Omega} \|W_{\theta}(x; \mu) - g(x; \mu)\|_{2}^{2} dx d\mu}_{\mathcal{J}_{\mathsf{boundary}}(\theta)}.$$

Both integrals are, once again, approximated by the Monte-Carlo method.
Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

- Why do we need well-balanced methods?
- Example of a physical model: the shallow water equations
- Numerical method overview: Discontinuous Galerkin
- Enhancing DG with Scientific Machine Learning
- Validation
- Conclusion

PINNs amount to solving a nonlinear optimization problem.

For such problems, state-of-the-art approaches rely on stochastic gradient descent⁴, and so require differentiating the loss function with respect to θ .

Because of the Monte-Carlo estimation, the loss function contains terms in $\mathcal{D}(W_{\theta}, x_j; \mu_i)$. Say \mathcal{D} contains a Laplace operator: we need to compute, among other things,

 $\nabla_{\theta} \Delta W_{\theta}(x_j; \mu_i).$

These differentials are exactly computed, thanks to automatic differentiation tools.

Fortunately, these tools are already implemented in several libraries (we used pytorch).

⁴Namely, on the ADAM algorithm: see D. Kingma and J. Ba, (2015).

PINNs are implemented in scimba⁵, developed in-house in the MACARON team.

The networks have 5 hidden layers of 20 neurons each, and $\sigma = \tanh$. In total, W_{θ} has 1761 parameters (one can compare this to a FEM with 1761 degrees of freedom). We train for 2 500 epochs (number of descent steps) and $N_c = 5000$ Monte-Carlo samples.

All computations are run on a single GPU, an AMD Instinct MI210.

We present PINN solutions, for several Ω and f, of a **four-dimensional parametric BVP**, whose solution depends on $x \in \Omega \subset \mathbb{R}^2$ and $\mu = (\alpha, \beta) \in \mathbb{M} \subset \mathbb{R}^2$:

$$\begin{cases} \Delta W(x;\mu) + \beta W(x;\mu) = f(x;\mu) & \text{ for } (x,\mu) \in \Omega \times \mathbb{M}, \\ W(x;\mu) = 0 & \text{ for } (x,\mu) \in \partial\Omega \times \mathbb{M}. \end{cases}$$

⁵freely accessible at https://gitlab.inria.fr/scimba/scimba

























Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

Context of hybrid methods

Advantages of the finite element method (FEM)

- The FEM is provably **convergent**: more DOFs lead to a more accurate solution.
- Optimized software is widely used in industry and academia.

Context of hybrid methods

Advantages of the finite element method (FEM)

- The FEM is provably **convergent**: more DOFs lead to a more accurate solution.
- Optimized software is widely used in industry and academia.

Advantages of PINNs

- PINNs are **mesh-less**, which is good for e.g. complex geometries.
- High-dimensional parametric problems are easily tackled.
- Once the network is trained, the solution inference is quick.

Context of hybrid methods

Advantages of the finite element method (FEM)

- The FEM is provably **convergent**: more DOFs lead to a more accurate solution.
- Optimized software is widely used in industry and academia.

Advantages of PINNs

- PINNs are **mesh-less**, which is good for e.g. complex geometries.
- High-dimensional parametric problems are easily tackled.
- Once the network is trained, the solution **inference is quick**.

Hybrid methods seek to combine the best of both worlds: in our case, using a PINN to improve the resolution of the FEM solution while retaining its order of accuracy.

We consider a parametric elliptic PDE $\mathcal{D}(u, x; \mu) = 0$.

We consider a parametric elliptic PDE $\mathcal{D}(u, x; \mu) = 0$.

We propose a two-step hybrid method:

- 1. **Offline phase**: train a neural network (e.g. a parametric PINN) to approximate a large family of solutions to the PDE;
- 2. **Online phase**: use the trained network to **correct the FEM approximation space**, and run the FEM simulation on a coarse grid.

Classical finite element method

The classical FEM relies on the following steps.

1. Rewrite the PDE $\mathcal{D}(u, x; \mu) = 0$ as a variational problem:

Find $u \in V$ such that $a(u, v) = \ell(v) \quad \forall v \in V$,

where $V = \mathcal{H}_0^m(\Omega)$ is a Hilbert space, *a* a bilinear form, and ℓ a linear form.

2. Discretize the domain Ω and introduce V_h a finite-dimensional subspace of V, to get

Find $u_h \in V_h$ such that $a(u_h, \mathbf{v}_h) = \ell(\mathbf{v}_h) \quad \forall \mathbf{v}_h \in V_h$.

3. Solve the above linear system to get the approximation u_h of u.

Classical finite element method

The classical FEM relies on the following steps.

1. Rewrite the PDE $\mathcal{D}(u, x; \mu) = 0$ as a variational problem:

Find $u \in V$ such that $a(u, v) = \ell(v) \quad \forall v \in V$,

where $V = \mathcal{H}_0^m(\Omega)$ is a Hilbert space, *a* a bilinear form, and ℓ a linear form.

2. Discretize the domain Ω and introduce V_h a finite-dimensional subspace of V, to get

Find $u_h \in V_h$ such that $a(u_h, \mathbf{v}_h) = \ell(\mathbf{v}_h) \quad \forall \mathbf{v}_h \in V_h$.

3. Solve the above linear system to get the approximation u_h of u.

The approximation space V_h is made of piecewise polynomial functions on the mesh \mathfrak{T}_h : $V_h = \{ \mathbf{v}_h \in \mathfrak{H}_0^m(\Omega) \cap C^0(\overline{\Omega}) \text{ such that } \forall K \in \mathfrak{T}_h, \mathbf{v}_h |_K \in \mathbb{P}_q(K) \}.$

Assume that we have a prior⁶ $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$ on the solution u.

 \rightsquigarrow How to use u_{θ} to **improve the FEM solution**?

⁶Here, given by a PINN, but that is not necessarily the case.

Assume that we have a prior⁶ $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$ on the solution u.

 \rightsquigarrow How to use u_{θ} to **improve the FEM solution**?

We suggest to modify the FEM approximation space, replacing V_h by V_h^+ , defined by:

 $V_h^+ = \{v_h = u_{\Theta} + p_h^+, \quad p_h^+ \in V_h\}.$

Since $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$, V_{h}^{+} remains a subspace of $\mathcal{H}_{0}^{m}(\Omega)$, like V_{h} .

⁶Here, given by a PINN, but that is not necessarily the case.

Assume that we have a prior⁶ $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$ on the solution u.

 \rightsquigarrow How to use u_{θ} to **improve the FEM solution**?

We suggest to modify the FEM approximation space, replacing V_h by V_h^+ , defined by:

$$V_h^+ = \{v_h = u_\theta + p_h^+, \quad p_h^+ \in V_h\}.$$

Since $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$, V_{h}^{+} remains a subspace of $\mathcal{H}_{0}^{m}(\Omega)$, like V_{h} .

The discrete variational problem becomes⁷:

$$\begin{pmatrix} \text{Find } u_h \in V_h \text{ such that} \\ \forall v_h \in V_h, \ a(u_h, v_h) = \ell(v_h) \end{pmatrix} \rightsquigarrow \begin{pmatrix} \text{Find } u_h^+ \in V_h^+ \text{ such that} \\ \forall v_h \in V_h, \ a(u_h, v_h) = \ell(v_h) \end{pmatrix}$$

⁶Here, given by a PINN, but that is not necessarily the case. ⁷This sets the method in the Petrov-Galerkin framework, where trial and test spaces are different.

Assume that we have a prior⁶ $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$ on the solution u.

 \rightsquigarrow How to use u_{θ} to **improve the FEM solution**?

We suggest to modify the FEM approximation space, replacing V_h by V_h^+ , defined by:

 $V_h^+ = \{v_h = u_\theta + p_h^+, \quad p_h^+ \in V_h\}.$

Since $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$, V_{h}^{+} remains a subspace of $\mathcal{H}_{0}^{m}(\Omega)$, like V_{h} .

The discrete variational problem becomes⁷:

 $\begin{pmatrix} \text{Find } u_h^+ \in V_h^+ \text{ such that} \\ \forall v_h \in V_h, \ a(u_h^+, v_h) = \ell(v_h) \end{pmatrix} \iff \begin{pmatrix} \text{Find } p_h^+ \in V_h \text{ such that} \\ \forall v_h \in V_h, \ a(p_h^+, v_h) = \ell(v_h) - a(u_\theta, v_h). \end{pmatrix}$

⁶Here, given by a PINN, but that is not necessarily the case.

⁷This sets the method in the Petrov-Galerkin framework, where trial and test spaces are different.

Error analysis (proof in appendix)

Equipped with the modified approximation space, we now perform an error analysis.

Theorem: Let $u \in \mathcal{H}_0^m$ be the exact solution of the BVP, $u_\theta \in \mathcal{H}_0^m(\Omega)$ a prior on u, and $u_h^+ \in V_h^+$ the enhanced FEM solution (considering \mathbb{P}_q polynomials, with $m \leq q$). Then:

$$\|u-u_h^+\|_{H^m} \lesssim C_{ ext{gain}}^+ \underbrace{h^{q+1-m}|u|_{H^{q+1}}}_{ ext{classical FEM error}}.$$

Error analysis (proof in appendix)

Equipped with the modified approximation space, we now perform an error analysis.

Theorem: Let $u \in \mathcal{H}_0^m$ be the exact solution of the BVP, $u_\theta \in \mathcal{H}_0^m(\Omega)$ a prior on u, and $u_h^+ \in V_h^+$ the enhanced FEM solution (considering \mathbb{P}_q polynomials, with $m \leq q$). Then:

$$\|u-u_h^+\|_{H^m}\lesssim C_{ ext{gain}}^+\underbrace{h^{q+1-m}|u|_{H^{q+1}}}_{ ext{classical FEM error}}.$$

In this result, the constant

$$C_{ ext{gain}}^+ = rac{|u - u_{\theta}|_{H^{q+1}}}{|u|_{H^{q+1}}}$$

represents the potential gain compared to the error of the classical FEM.

Key remark: The prior u_{θ} must be a good approximation of the $(q + 1)^{\text{th}}$ derivative of u. This is why we use PINNs, rather than purely data-driven priors! This hybrid method can be seen as

- enhancing the FEM approximation space with a PINN prior, to get V_h^+ ;
- or ensuring the convergence of a PINN approximation by using a coarse FEM.

Remark: The hybrid method consists in offline and online parts:Offline: Train the PINN on the parametric PDE (potentially time-consuming).Online: There are two online substeps:

evaluate the NN at Gauss points to compute the approximation space,
use a regular, coarse FEM solver with the new approximation space.
NN inference is quick, so the online cost of using the NN is negligible!

4D Poisson problem

First, we tackle the following 4D PDE (2D in space, with two parameters):

$$\begin{cases} -\Delta u = f & \text{in } \Omega, \\ u = 0 & \text{on } \partial \Omega, \end{cases}$$

with $\Omega = (-\frac{\pi}{2}, \frac{\pi}{2})^2$, parameters $x_1^0, x_2^0 \sim \mathcal{U}(-0.5, 0.5)$ and exact solution

$$u(x_1, x_2; x_1^0, x_2^0) = \sin(2x_1)\sin(2x_2)\exp\left(-\frac{1}{2}\left((x_1 - x_1^0)^2 + (x_2 - x_2^0)^2\right)\right).$$

4D Poisson problem

First, we tackle the following 4D PDE (2D in space, with two parameters):

$$\begin{cases} -\Delta u = f & \text{in } \Omega, \\ u = 0 & \text{on } \partial \Omega, \end{cases}$$

with $\Omega=(-\frac{\pi}{2},\frac{\pi}{2})^2$, parameters $x_1^0,x_2^0\sim \mathcal{U}(-0.5,0.5)$ and exact solution

$$u(x_1, x_2; x_1^0, x_2^0) = \sin(2x_1)\sin(2x_2)\exp\left(-\frac{1}{2}\left((x_1 - x_1^0)^2 + (x_2 - x_2^0)^2\right)\right).$$

With a given q, we compare, averaging over 50 values of the parameters (x_1^0, x_2^0) , the relative L^2 errors of the enhanced \mathbb{P}_q FEM (with approximation space V_h^+) to

- the classical \mathbb{P}_q FEM (with approximation space V_h);
- the results of the PINN.

4D Poisson problem – gains

We add a component to the loss function: the derivatives with respect to the parameters

 $\|\partial_{x_1^0}(\Delta u_\theta+f)\|+\|\partial_{x_2^0}(\Delta u_\theta+f)\|.$

4D Poisson problem – gains

We add a component to the loss function: the derivatives with respect to the parameters

 $\|\partial_{x_1^0}(\Delta u_\theta+f)\|+\|\partial_{x_2^0}(\Delta u_\theta+f)\|.$

		Gains (L ² error): ours w.r.t. PINNs			Gains (L ² error): ours w.r.t. FEM		
q	Ν	min	max	mean	min	max	mean
1	20	18.28	66.19	43.42	243.79	874.3	633.45
1	40	73.45	272.36	176.52	241.8	843.29	621.68
2	20	362.57	2,052.78	1,025.28	177.74	476.76	376.16
2	40	3,081.22	17,532.62	8,725.57	177.16	472.55	371.93
3	20	4,879.13	32,757.68	14,646.89	116.52	298.33	208.35
3	40	88,736.63	587,716.86	264,383.45	117.46	296.34	208.29



4D Poisson problem – computation time

We now compare computation times: we record the mesh size and the computation time *T* (excluding training time! see next slide) required to reach an error *E*.


4D Poisson problem – training time and parametric PDEs

We study the cost-effectiveness of our method in for parametric PDEs. Namely, we **run** n_p **simulations with** n_p **different parameter sets**.

The total time to perform n_p simulations is the following:

for V_h : $T_{V_h}^{\text{total}} = n_p \times T_{V_h}$; for V_h^+ : $T_{V_h^+}^{\text{total}} = T_{\text{training}} + n_p \times T_{V_h^+}$.

To reach an error of 10^{-3} , $T_{V_h} \simeq 43s$ and $T_{V_h^+} \simeq 0.24s$.

We study the cost-effectiveness of our method in for parametric PDEs. Namely, we **run** n_p **simulations with** n_p **different parameter sets**.

The total time to perform n_p simulations is the following:

for V_h : $T_{V_h}^{\text{total}} = n_p \times T_{V_h}$; for V_h^+ : $T_{V_h^+}^{\text{total}} = T_{\text{training}} + n_p \times T_{V_h^+}$.

To reach an error of 10^{-3} , $T_{V_h} \simeq 43s$ and $T_{V_h^+} \simeq 0.24s$.

Therefore, with a training time $T_{\text{training}} \simeq 240$ s, **our method is cost-effective for** $n_p \ge 6$.

This figure also depends on q and on the error one wishes to reach for each simulation. Also, our FEM code is not highly optimized; with a faster code, n_p would be larger.

4D Poisson problem on a donut

We now consider the Poisson problem on a donut, with Dirichlet boundary conditions.



		Gains (L ² error): ours w.r.t. PINNs			Gains (L ² error): ours w.r.t. FEM		
q	Ν	min	max	mean	min	max	mean
1	20	10.18	35.8	19.49	71.17	254.32	153.44
1	40	33.35	125.03	65.64	63.93	199.95	131.06
2	20	189.1	1,331.27	485.95	32.47	80.69	58.98
2	40	1,241.42	9,686.46	3,261.71	30.57	74.15	54.09
3	20	5,630.17	39,651.58	14,987.25	15.73	32.1	23.07
3	40	74,794.74	573,631.63	202,631.9	13.67	29.52	20.57

Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

Well-balanced DG methods using bases enriched with PINNs Why do we need well-balanced methods?

Tsunami simulation: naive numerical method

Tsunami initialization

Simulation with a naive numerical method



Tsunami simulation: naive numerical method

Tsunami initialization



Simulation with a naive numerical method



~> The simulation is not usable!

Indeed, the ocean at rest, far from the tsunami, started spontaneously producing waves.

This comes from the non-preservation of stationary solutions, hence the need to develop numerical methods that **preserve stationary solutions**: so-called **well-balanced** methods.

Tsunami simulation: well-balanced method



Victor Michel-Dansac | Hybrid methods for elliptic and hyperbolic PDEs | Workshop MEDIA, La Garde

Tsunami simulation: well-balanced method



Victor Michel-Dansac | Hybrid methods for elliptic and hyperbolic PDEs | Workshop MEDIA, La Garde

The goal of this work is to provide a numerical method which:

- is able to deal with generic systems of balance laws,
- can provide a very good approximation of families of steady solutions,
- is as accurate as classical methods on unsteady solutions,
- with provable convergence estimates.

To that end, we select the **Discontinuous Galerkin (DG)** framework.

Well-balanced DG methods using bases enriched with PINNs Example of a physical model: the shallow water equations

The shallow water equations

The shallow water equations are governed by the following PDE:

$$\begin{cases} \partial_t h + \partial_x q = 0, \\ \partial_t q + \partial_x \left(\frac{q^2}{h} + \frac{1}{2}gh^2 \right) = -gh\partial_x Z(x). \end{cases}$$



- h(x, t): water depth
- u(x, t): water velocity
- q = hu: water discharge
- Z(x): known topography
- g: gravity constant

The shallow water equations: steady solutions

The steady solutions of the shallow water equations are governed by the following ODEs:

$$\begin{cases} \partial_x q = 0, \\ \partial_x \left(\frac{q^2}{h} + \frac{1}{2}gh^2 \right) = -gh\partial_x Z(x), \end{cases} \xrightarrow{\sim}$$

$$\begin{cases} q = \operatorname{cst} \rightleftharpoons q_0, \\ \frac{q_0^2}{2h^2} + g(h+Z) = \operatorname{cst.} \end{cases}$$



If the velocity vanishes, i.e. $q_0 = 0$, we obtain the lake at rest steady solution:

 $h+Z=\operatorname{cst}=H_0.$

The shallow water equations: steady solutions

The steady solutions of the shallow water equations are governed by the following ODEs:

 \rightarrow

$$\begin{cases} \partial_x q = 0, \\ \partial_x \left(\frac{q^2}{h} + \frac{1}{2}gh^2 \right) = -gh\partial_x Z(x), \end{cases}$$

$$\begin{cases} q = \operatorname{cst} \rightleftharpoons q_0, \\ \frac{q_0^2}{2h^2} + g(h+Z) = \operatorname{cst.} \end{cases}$$



For a nonzero discharge $q_0 \neq 0$, we obtain a moving steady solution: h(x) satisfies a polynomial equation of degree 3 for all x.

Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

Finite volume method, visualized



Discontinuous Galerkin, visualized



Discontinuous Galerkin, visualized



On the previous slide, the data *W* is represented by

- a polynomial of degree 2 in each cell (Galerkin approximation),
- which is **Discontinuous** at interfaces between cells.

On the previous slide, the data *W* is represented by

- a polynomial of degree 2 in each cell (Galerkin approximation),
- which is **Discontinuous** at interfaces between cells.

Therefore, in each cell Ω_i , *W* is approximated by

$$\left. \mathsf{W} \right|_{\Omega_i} \simeq \mathsf{W}_i^{ extsf{DG}} \coloneqq lpha_0 + lpha_1 \mathsf{X} + lpha_2 \mathsf{X}^2 = \sum_{j=0}^2 lpha_j \mathsf{X}^j,$$

where the polynomial coefficients α_0 , α_1 and α_2 are determined to ensure fitness between the continuous data and its polynomial approximation.

Any polynomial of degree two can be exactly represented this way.

More generally, we define a polynomial basis $\varphi_0, \ldots, \varphi_N$ on each cell Ω_i and approximate the solution in this basis.

A usual example is the following so-called **modal basis**:

$$\forall j \in \{0,\ldots,N\}, \quad \varphi_j(x) = x^j.$$

More generally, we define a polynomial basis $\varphi_0, \ldots, \varphi_N$ on each cell Ω_i and approximate the solution in this basis.

A usual example is the following so-called **modal basis**:

$$\forall j \in \{0,\ldots,N\}, \quad \varphi_j(x) = x^j.$$

Main takeaway: The DG scheme is exact on every function that can be exactly represented in the basis!

Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

Recall that the DG scheme will be exact on every function that can be exactly represented in the DG basis, as soon as it is also a solution to the PDE.

Recall that the DG scheme will be exact on every function that can be exactly represented in the DG basis, as soon as it is also a solution to the PDE.

Main idea

Enhance the DG basis by using the steady solution!

---- If the steady solution or an approximation thereof is contained in the basis, then:

- using the exact steady solution in the basis will make the scheme exactly wellbalanced;
- using an approximation of the steady solution will make the scheme approximately well-balanced.

Enhanced DG bases

Assume that you know a **prior** W_{θ} on the steady solution.

It can be the exact steady solution ($W_{\theta} = W_{eq}$), or it can be an approximation ($W_{\theta} \simeq W_{eq}$).

The goal is now to **enhance the modal basis** V using W_{θ} :

$$V = \{1, x, x^2, \ldots, x^N\}.$$

Enhanced DG bases

Assume that you know a **prior** W_{θ} on the steady solution.

It can be the exact steady solution ($W_{\theta} = W_{eq}$), or it can be an approximation ($W_{\theta} \simeq W_{eq}$).

The goal is now to **enhance the modal basis** V using W_{θ} :

$$V = \{1, x, x^2, \ldots, x^N\}.$$

First possibility: multiply the whole basis by W_{θ}

$$V^{\theta}_* = \{W_{\theta}, x W_{\theta}, x^2 W_{\theta}, \dots, x^N W_{\theta}\}.$$

Enhanced DG bases

Assume that you know a **prior** W_{θ} on the steady solution.

It can be the exact steady solution ($W_{\theta} = W_{eq}$), or it can be an approximation ($W_{\theta} \simeq W_{eq}$).

The goal is now to **enhance the modal basis** V using W_{θ} :

$$V = \{1, x, x^2, \ldots, x^N\}.$$

First possibility: multiply the whole basis by W_{θ}

$$V^{\theta}_* = \{W_{\theta}, x W_{\theta}, x^2 W_{\theta}, \dots, x^N W_{\theta}\}.$$

Second possibility: replace the first element with W_{θ}

$$V^{\theta}_{+} = \{ W_{\theta}, x, x^2, \dots, x^N \}.$$

Error estimates

We denote by:

- W_{ex} the exact solution,
- W_{DG} the approximate solution without prior,
- $W_{\rm DG}^{\theta}$ the approximate solution with prior W_{θ} and basis V_{*}^{θ} .

For a DG scheme of order q + 1, we obtain⁸ the following error estimates:

$$\begin{split} \|W_{\mathrm{ex}} - W_{\mathrm{DG}}\| &\lesssim \left|W_{\mathrm{ex}}\right|_{H^{q+1}} \Delta x^{q+1}, \\ \|W_{\mathrm{ex}} - W_{\mathrm{DG}}^{\theta}\| &\lesssim \left|\frac{W_{\mathrm{ex}}}{W_{\theta}}\right|_{H^{q+1}} \Delta x^{q+1} \|W_{\theta}\|_{L^{\infty}}. \end{split}$$

Conclusion of the error estimates: the prior W_{θ} needs to provide a **good approximation of the derivatives** of the steady solution.

⁸Rigorous error estimates are written in terms of the error in the projection onto both bases.

For very simple systems, one can use the exact steady solution as a prior.

However, in many cases, even for some simple and well-known systems, one cannot compute the exact steady solution. Therefore, **an approximation is required**.

How to obtain such an approximation?

For very simple systems, one can use the exact steady solution as a prior.

However, in many cases, even for some simple and well-known systems, one cannot compute the exact steady solution. Therefore, **an approximation is required**.

How to obtain such an approximation?

- 1. **First possibility**: use a traditional numerical approximation, obtained by classical ODE solvers (e.g. Runge-Kutta schemes).
- 2. Second possibility: use a Physics-Informed Neural Network (PINN).

Since we need a good approximation of the derivatives, we use a PINN.

Next step: Validate the method with several numerical experiments.

Introduction – framework for approximating solutions to PDEs

Physics-Informed Neural Networks (PINNs)

Using PINNs for parametric PDEs

Numerical examples

Hybridizing the finite element method and PINNs

Well-balanced DG methods using bases enriched with PINNs

Why do we need well-balanced methods?

Example of a physical model: the shallow water equations

Numerical method overview: Discontinuous Galerkin

Enhancing DG with Scientific Machine Learning

Validation

Conclusion

Setup: the advection equation

We run experiments on the **advection equation with source term**, with a given initial condition $W_0 : \mathbb{R} \to \mathbb{R}$:

$$\begin{cases} \partial_t W + c \partial_x W = a W + b W^2 & \text{ for } x \in (0, 1), \ t \in (0, T), \\ W(0, x) = W_0(x) & \text{ for } x \in (0, 1), \\ W(t, 0) = u_0 & \text{ for } t \in (0, T). \end{cases}$$

The steady solution W_{eq} satisfies the BVP

$$\begin{cases} c\partial_x W_{\text{eq}} - aW_{\text{eq}} - bW_{\text{eq}}^2 = 0 \quad \text{for } x \in (0, 1), \\ W_{\text{eq}}(0) = u_0, \end{cases}$$

whose unique solution is, with parameters $\mu = \{a, b, c, u_0\}$:

$$W_{\text{eq}}(x;\mu) = \frac{au_0}{(a+bu_0)e^{-\frac{ax}{c}}-bu_0}.$$
Thanks to the boundary ansatz and the ODE loss, the final loss function **does not need any data**, and there is **no competition between loss functions**: we get

$$\mathcal{J}(\theta) = \int_{\mathbb{P}} \int_{\Omega} \left\| c \partial_x \widetilde{W_{\theta}} - a \widetilde{W_{\theta}} - b \widetilde{W_{\theta}}^2 \right\|_2^2 dx d\mu,$$

with the ansatz

$$\widetilde{W_{\theta}}=u_{0}+x\,W_{\theta},$$

with W_{θ} the result of the neural network.

In practice, we take c = 1 and make sure the steady solution is well-defined, by taking

$$\mathbb{P} = \{(a, b, u_0) \in (0.5, 1) \times (0.5, 1) \times (0.1, 0.2)\}.$$

Hence, the neural network is a function $W_{\theta} \in \mathcal{C}^{\infty}(\mathbb{R} \times \mathbb{R}^3, \mathbb{R})$.

We use the DG scheme to solve the advection equation with the **steady solution as initial condition**:

 $\begin{cases} \partial_t W + \partial_x W = aW + bW^2 & \text{ for } x \in (0, 1), \ t \in (0, T), \\ W(0, x) = W_{eq}(x) & \text{ for } x \in (0, 1), \\ W(t, 0) = u_0 & \text{ for } t \in (0, T). \end{cases}$

We expect the DG scheme with prior:

- to provide a **better approximation of the steady solution** than the classical DG scheme (approximate well-balanced property),
- while converging with the same order of accuracy.

We compute the errors in x between the exact and approximate solutions:

• for several numbers of basis elements and discretization cells,

• using
$$a = 0.75$$
; $b = 0.75$; $u_0 = 0.15$.

	without prior			W		
cells	error	order		error	order	gain
10	$1.75 \cdot 10^{-2}$	_		$1.45 \cdot 10^{-5}$	_	1200.02
20	$8.75 \cdot 10^{-3}$	1.00		$7.61 \cdot 10^{-6}$	0.93	1149.11
40	$4.38 \cdot 10^{-3}$	1.00		$3.92 \cdot 10^{-6}$	0.96	1118.29
80	$2.19 \cdot 10^{-3}$	1.00		$2.00\cdot10^{-6}$	0.97	1098.77
160	$1.10 \cdot 10^{-3}$	1.00		$1.01 \cdot 10^{-6}$	0.98	1085.96

(a) Errors with a basis composed of one element.

We compute the errors in x between the exact and approximate solutions:

• for several numbers of basis elements and discretization cells,

• using
$$a = 0.75$$
; $b = 0.75$; $u_0 = 0.15$.

	without prior			with prior			
cells	error	order		error	order	gain	
10	$4.93 \cdot 10^{-4}$	_		$2.18 \cdot 10^{-6}$	_	226.00	
20	$1.24 \cdot 10^{-4}$	2.00		$3.20 \cdot 10^{-7}$	2.77	386.66	
40	$3.09 \cdot 10^{-5}$	2.00		$8.07 \cdot 10^{-8}$	1.99	382.88	
80	$7.72 \cdot 10^{-6}$	2.00		$2.05 \cdot 10^{-8}$	1.98	376.52	
160	$1.93 \cdot 10^{-6}$	2.00		$5.16 \cdot 10^{-9}$	1.99	374.49	

(b) Errors with a basis composed of two elements.

We compute the errors in x between the exact and approximate solutions:

• for several numbers of basis elements and discretization cells,

• using
$$a = 0.75$$
; $b = 0.75$; $u_0 = 0.15$.

	without prior			with prior			
cells	error	order		error	order	gain	
10	7.89 · 10 ⁻⁶	_		$1.00 \cdot 10^{-7}$	_	78.58	
20	$9.94 \cdot 10^{-7}$	2.99		$1.33 \cdot 10^{-8}$	2.91	74.60	
40	$1.24 \cdot 10^{-7}$	3.00		$1.72 \cdot 10^{-9}$	2.95	72.13	
80	$1.55 \cdot 10^{-8}$	3.00		$2.17 \cdot 10^{-10}$	2.99	71.43	
160	$1.94 \cdot 10^{-9}$	3.00		$2.72 \cdot 10^{-11}$	3.00	71.25	

(c) Errors with a basis composed of three elements.

We compute the errors in x between the exact and approximate solutions:

• for several numbers of basis elements and discretization cells,

• using
$$a = 0.75$$
; $b = 0.75$; $u_0 = 0.15$.

	without p	without prior			with prior			
cells	error	order		error	order	gain		
10	$1.20 \cdot 10^{-7}$	_		8.31 · 10 ⁻⁹	_	14.40		
20	$7.39 \cdot 10^{-9}$	4.02		$5.51 \cdot 10^{-10}$	3.91	13.40		
40	$4.59 \cdot 10^{-10}$	4.01		$3.48 \cdot 10^{-11}$	3.99	13.19		
80	$2.92 \cdot 10^{-11}$	3.98		$2.20 \cdot 10^{-12}$	3.99	13.27		
160	$1.85 \cdot 10^{-12}$	3.98		$1.29 \cdot 10^{-13}$	4.10	14.38		

(d) Errors with a basis composed of four elements.

To study the influence of the parameters, we compute some statistics on the gains with 1000 random sets of parameters in \mathbb{P} , for a DG scheme of order q + 1.

q	minimum gain	average gain	maximum gain
0	63.46	735.08	4571.89
1	32.22	149.38	450.74
2	6.20	54.16	118.45
3	1.55	19.54	108.10

We also compare the computation time in bases V and V^{θ}_{+} :

- **assembly** time records meshing the domain and assembling the DG matrices (which requires evaluating the prior);
- **scheme** time records performing a full time loop of the DG scheme.

Computation times are reported in seconds.

	without prior			with prior			
cells	assembly	scheme	total	assembly	scheme	total	ratio
10	7.08e-4	1.98e-2	2.06e-2	5.98e-3	1.98e-2	2.58e-2	1.26
20	6.78e-4	3.77e-2	3.84e-2	5.81e-3	3.78e-2	4.36e-2	1.14
40	7.18e-4	7.70e-2	7.77e-2	6.04e-3	7.70e-2	8.30e-2	1.07
80	7.41e-4	1.57e-1	1.57e-1	6.24e-3	1.57e-1	1.63e-1	1.04
160	7.61e-4	3.16e-1	3.17e-1	6.40e-3	3.13e-1	3.20e-1	1.01

(e) Computation time with a basis composed of one element.

We also compare the computation time in bases V and V^{θ}_+ :

- **assembly** time records meshing the domain and assembling the DG matrices (which requires evaluating the prior);
- **scheme** time records performing a full time loop of the DG scheme.

Computation times are reported in seconds.

	without prior			with prior			
cells	assembly	scheme	total	assembly	scheme	total	ratio
10	7.45e-4	4.07e-2	4.14e-2	5.81e-3	4.19e-2	4.78e-2	1.15
20	7.52e-4	8.19e-2	8.27e-2	5.94e-3	8.15e-2	8.74e-2	1.06
40	8.09e-4	1.72e-1	1.73e-1	6.09e-3	1.68e-1	1.74e-1	1.01
80	8.43e-4	3.52e-1	3.53e-1	6.12e-3	3.50e-1	3.56e-1	1.01
160	1.00e-3	7.45e-1	7.46e-1	6.50e-3	7.44e-1	7.50e-1	1.00

(f) Computation time with a basis composed of two elements.

We also compare the computation time in bases V and V^{θ}_{+} :

- **assembly** time records meshing the domain and assembling the DG matrices (which requires evaluating the prior);
- **scheme** time records performing a full time loop of the DG scheme.

Computation times are reported in seconds.

	without prior			with prior			
cells	assembly	scheme	total	assembly	scheme	total	ratio
10	8.86e-4	4.04e-2	4.13e-2	6.17e-3	4.10e-2	4.72e-2	1.14
20	9.28e-4	8.21e-2	8.31e-2	6.14e-3	8.41e-2	9.03e-2	1.09
40	9.84e-4	1.75e-1	1.76e-1	6.32e-3	1.75e-1	1.81e-1	1.03
80	1.04e-3	3.60e-1	3.61e-1	6.39e-3	3.63e-1	3.69e-1	1.02
160	1.15e-3	7.95e-1	7.96e-1	6.94e-3	7.98e-1	8.05e-1	1.01

(g) Computation time with a basis composed of three elements.

We also compare the computation time in bases V and V^{θ}_+ :

- **assembly** time records meshing the domain and assembling the DG matrices (which requires evaluating the prior);
- **scheme** time records performing a full time loop of the DG scheme.

Computation times are reported in seconds.

	without prior			with prior			
cells	assembly	scheme	total	assembly	scheme	total	ratio
10	1.02e-3	7.25e-2	7.35e-2	6.31e-3	7.25e-2	7.88e-2	1.07
20	1.03e-3	1.47e-1	1.48e-1	6.35e-3	1.47e-1	1.53e-1	1.04
40	1.11e-3	3.07e-1	3.08e-1	6.49e-3	3.00e-1	3.06e-1	1.00
80	1.24e-3	6.47e-1	6.48e-1	6.67e-3	6.50e-1	6.56e-1	1.01
160	1.34e-3	1.49	1.49	7.12e-3	1.50	1.50	1.01

(h) Computation time with a basis composed of four elements.

Perturbation of a shallow water steady solution



PINN trained on a parametric steady solution, driven by the topography

 $Z(x; \mu) = \Gamma \exp\left(\alpha (r_0^2 - \|x\|^2)\right),$

with physical parameters

μ

$$\in \mathbb{P} \iff egin{cases} lpha \in [0.25, 0.75], \ \Gamma \in [0.1, 0.4], \ r_0 \in [0.5, 1.25]. \end{cases}$$

Left plot: initial condition, made of a perturbed steady solution.

Perturbation of a shallow water steady solution



Conclusion

Conclusion

We introduced

- a framework for approximating solutions to PDEs with linear or nonlinear functions,
- physics-informed neural networks (PINNs),
- a hybrid method between FEM and PINNs, applied to elliptic problems,
- a hybrid method blending physics-informed learning and DG bases.

Perspectives include

- · tackling time-dependent solutions,
- going to complex three-dimensional geometries and richer PDEs.

Related paper: E. Franck, V. Michel-Dansac and L. Navoret.

"Approximately WB DG methods using bases enriched with PINNs.", J. Comput. Phys., 2024 git repository: https://github.com/Victor-MichelDansac/DG-PINNs

Thank you for your attention!

For the moment, the **boundary conditions are viewed as constraints**, and the solution will not exactly satisfy them.

This can be remedied by introducing a **suitable ansatz**⁹. To that end, we define

 $\widetilde{W_\theta}=\mathfrak{B}(W_\theta,x,t;\mu),\quad\text{such that}\quad \widetilde{W_\theta}(x,t;\mu)=g(x,t;\mu)\quad\text{for }x\in\partial\Omega.$

Clearly, the new approximate solution $\widetilde{W_{\theta}}$ exactly satifies the boundary conditions.

Moreover, the boundary loss function can be eliminated, thus **reducing competition** between the loss functions.

 \rightsquigarrow How to get such an ansatz? We check on an example.

⁹I. E. Lagaris et al., IEEE Trans. Neural Netw. (1998)

Exact imposition of the boundary conditions: example

Let us go back to the parameterized Laplace equation, where $\mu=(\alpha,\beta)$:

$$\begin{cases} \Delta W(x; \mu) + \beta W(x; \mu) = f(x; \mu) & \text{ for } (x, \mu) \in \Omega \times \mathbb{P}, \\ W(x; \mu) = 0 & \text{ for } (x, \mu) \in \partial\Omega \times \mathbb{P}. \end{cases}$$

Homogeneous Dirichlet BC are imposed on $\Omega = (0,1)^2$, and so we define the ansatz

$$\widetilde{W_{\theta}} = \mathcal{B}(W_{\theta}, \mathbf{x}; \boldsymbol{\mu}) = x_1(1 - x_1) x_2(1 - x_2) W_{\theta}.$$

This obviously satisfies the boundary conditions, since $\forall x \in \partial \Omega, \widetilde{W_{\theta}}(x; \mu) = 0$.

Therefore, the loss function only has to ensure that $\widetilde{W_{\theta}}$ approximates the solution to the PDE in the interior of Ω , through minimizing the loss function

$$\mathcal{J}_{\mathsf{PDE}}(\theta) = \int_{\mathbb{P}} \int_{\Omega} \left\| \Delta \widetilde{W_{\theta}}(x; \mu) + \beta \widetilde{W_{\theta}}(x; \mu) - f(x; \mu) \right\|_{2}^{2} dx \, d\mu.$$

Spectral bias: MLPs first learn the low frequencies, before learning the high ones (with difficulty).

To illustrate this, we consider the high-frequency solution

$$W_{\text{exact}}(x;\mu) = \alpha\beta\sin(8\pi x_1)\sin(8\pi x_2).$$











To overcome the spectral bias of MLPs, we can use Fourier features¹⁰.

In this case, we replace the call to the neural network, going from $W_{\theta}(x; \mu)$ to

 $W_{\theta}(x; \mu, \sin(\pi a_1 x), \cos(\pi b_1 x), \dots, \sin(\pi a_K x), \cos(\pi b_K x)),$

with $K \in \mathbb{N}$ the number of Fourier features and $(a_i)_i$, $(b_i)_i$ the trainable frequencies.

¹⁰See [M. Tancik et al, (2021)], but other methods exist, such as Finite Basis PINNs (FBPINNs, see [V. Dolean et al., *Comput. Method. Appl. M.* (2024)]).



elapsed time: $\sim 50~seconds$







Monte-Carlo integration: convergence

Consider an integrable and bounded function $f : \Omega \times (0,T) \to \mathbb{R}$, and define $(X_k, T_k)_k$ a sequence of independent random variables, uniformly sampled in $\Omega \times (0,T)$.

We wish to give an approximation to

$$I=\int_{\Omega}\int_0^T f(x,t)\,dt\,dx.$$

An estimator of *I* is the following:

$$\widehat{m} = \frac{|\Omega|T}{n} \sum_{k=1}^{n} f(X_k, T_k).$$

Since the $(X_k, T_k)_k$ are uniformly sampled, we get

$$\mathbb{E}[f(X_k,T_k)] = \frac{1}{|\Omega|T} \int_{\Omega} \int_0^T f(x,t) \, dt \, dx.$$

Hence, applying the **law of large numbers** tells us that, with probability 1, $\hat{I^n} \rightarrow I$.

We can also determine the convergence speed of the Monte-Carlo method, assuming that f^2 is integrable.

The central limit theorem allows us to state that

$$\sqrt{n}(\widehat{I^n}-I)\xrightarrow[n\to\infty]{} \mathcal{N}(\mathbf{0},\sigma^2),$$

and so, for large enough *n* and with probability 1,

 $\left|\widehat{I^{n}}-I\right|=\mathfrak{O}(n^{-\frac{1}{2}}).$

This result is **independent of the dimension** d of Ω ! Contrast this with, for instance, the trapezoidal rule, with an error in $O(n^{-\frac{2}{d}})$.

To prove this result, we adapt the proof of Céa's lemma to the additive prior case. The numerical solution u_h^+ is given for all $x \in \Omega$ by

 $u_h^+(x) = u_\theta(x) + p_h^+(x),$

with $p_h^+ \in V_h \subset V$ solution of the new discrete variational problem. We have

$$\begin{aligned} a(u - u_{h}^{+}, u - u_{h}^{+}) &= a(u - u_{h}^{+}, (u - u_{\theta}) - p_{h}^{+}) \\ &= a(u - u_{h}^{+}, (u - u_{\theta}) - p_{h}^{+} - v_{h} + v_{h}), \qquad \forall v_{h} \in V_{h} \\ &= a(u - u_{h}^{+}, (u - u_{\theta}) - v_{h}) + a(u - u_{h}^{+}, v_{h} - p_{h}^{+}), \quad \forall v_{h} \in V_{h}. \end{aligned}$$

We will estimate both terms, one by one.

Let us first estimate the second term: $a(u - u_h^+, v_h - p_h^+)$.

Using that $V_h \subset V$, we have, by Galerkin orthogonality,

$$a(u-u_h^+,v_h)=0, \quad \forall v_h \in V_h.$$

The above equality is valid for all $v_h \in V_h$, and $v_h - p_h^+ \in V_h$. Therefore, we obtain

$$a(u-u_h^+,v_h-p_h^+)=0, \quad \forall v_h \in V_h.$$

The second term therefore vanishes, and we are left with the first one:

$$a(u-u_h^+,u-u_h^+)=a(u-u_h^+,(u-u_\theta)-v_h), \quad \forall v_h \in V_h.$$

Error analysis - proof (cont'd)

Denoting by α and γ the coercivity and continuity constants of *a*, we have

$$\begin{split} \alpha \big\| u - u_h^+ \big\|_{H^m}^2 &\leqslant a \big(u - u_h^+, u - u_h^+ \big) = a \big(u - u_h^+, (u - u_\theta) - v_h \big), &\forall v_h \in V_h, \\ &\leqslant \gamma \big\| u - u_h^+ \big\|_{H^m} \big\| (u - u_\theta) - v_h \big\|_{H^m}, &\forall v_h \in V_h, \end{split}$$

which immediately leads to

$$\|u-u_h^+\|_{H^m} \leqslant \frac{\gamma}{\alpha} \|(u-u_\theta)-v_h\|_{H^m}, \quad \forall v_h \in V_h.$$

Applying the above relation to $v_h = \mathcal{I}_h(u - u_\theta)$ with \mathcal{I}_h the Lagrange interpolator, and invoking classical interpolation results from [A. Ern and J.-L. Guermond, (2004)], we get

$$\|u-u_h^+\|_{H^m}\lesssim \frac{\gamma}{\alpha}h^{q+1-m}|u-u_\theta|_{H^{q+1}}.$$

Rewriting the above equation to introduce the error of the classical FEM, we get

$$\|u - u_h^+\|_{H^m} \lesssim C_{ ext{gain}} h^{q+1-m} |u|_{H^{q+1}} \quad ext{with} \quad C_{ ext{gain}} = rac{|u - u_{ ext{θ}}|_{H^{q+1}}}{|u|_{H^{q+1}}},$$

which completes the proof.

Enhancing the approximation space – multiplicative prior

Another possible modification of the FEM approximation space, is to replace V_h by V_h^{\times} :

$$V_h^{ imes} = ig\{ v_h = u_{ heta} p_h^{ imes}, \quad p_h^{ imes} \in V_h ig\}.$$

The discrete variational problem becomes:

$$\begin{pmatrix} \text{Find } u_h^{\times} \in V_h^{\times} \text{ such that} \\ \forall v_h \in V_h, \ a(u_h, v_h) = \ell(v_h) \end{pmatrix} \iff \begin{pmatrix} \text{Find } p_h \in V_h \text{ such that} \\ \forall v_h \in V_h, \ a(u_0 p_h, v_h) = \ell(v_h) \end{pmatrix}$$

Enhancing the approximation space – multiplicative prior

Another possible modification of the FEM approximation space, is to replace V_h by V_h^{\times} :

$$V_h^{ imes} = ig\{ v_h = u_{ heta} p_h^{ imes}, \quad p_h^{ imes} \in V_h ig\}.$$

The discrete variational problem becomes:

$$\begin{pmatrix} \text{Find } u_h^{\times} \in V_h^{\times} \text{ such that} \\ \forall v_h \in V_h, \ a(u_h, v_h) = \ell(v_h) \end{pmatrix} \iff \begin{pmatrix} \text{Find } p_h \in V_h \text{ such that} \\ \forall v_h \in V_h, \ a(u_\theta p_h, v_h) = \ell(v_h) \end{pmatrix}$$

Theorem: Let *u* be the exact solution of the BVP, $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$ a prior on *u*, and $u_{h}^{\times} \in V_{h}^{\times}$ the enhanced FEM solution (considering \mathbb{P}_{q} polynomials, with $m \leq q$). Then:

$$\|u-u_h^{\times}\|_{H^m} \lesssim C_{\mathrm{gain}}^{\times} h^{q+1-m} |u|_{H^{q+1}}.$$
Enhancing the approximation space – multiplicative prior

Another possible modification of the FEM approximation space, is to replace V_h by V_h^{\times} :

$$V_h^{ imes} = ig\{ v_h = u_{ heta} p_h^{ imes}, \quad p_h^{ imes} \in V_h ig\}.$$

The discrete variational problem becomes:

$$\begin{pmatrix} \text{Find } u_h^{\times} \in V_h^{\times} \text{ such that} \\ \forall v_h \in V_h, \ a(u_h, v_h) = \ell(v_h) \end{pmatrix} \iff \begin{pmatrix} \text{Find } p_h \in V_h \text{ such that} \\ \forall v_h \in V_h, \ a(u_\theta p_h, v_h) = \ell(v_h). \end{pmatrix}$$

Theorem: Let *u* be the exact solution of the BVP, $u_{\theta} \in \mathcal{H}_{0}^{m}(\Omega)$ a prior on *u*, and $u_{h}^{\times} \in V_{h}^{\times}$ the enhanced FEM solution (considering \mathbb{P}_{q} polynomials, with $m \leq q$). Then:

$$\|u-u_h^{\times}\|_{H^m} \lesssim C_{\mathrm{gain}}^{\times} h^{q+1-m} |u|_{H^{q+1}}.$$

In this result, the gain constant is $C_{\text{gain}}^{\times} = \left| \frac{u}{u_{\theta}} \right|_{H^{q+1}} \frac{\|u_{\theta}\|_{W^{m,\infty}}}{|u|_{H^{q+1}}}$. Beware of the division!

We use the DG scheme to solve the advection equation with a **perturbation of the steady solution as initial condition**:

 $\begin{cases} \partial_t W + \partial_x W = aW + bW^2 & \text{for } x \in (0,1), \ t \in (0,T), \\ W(0,x) = (1 + \varepsilon \sin(2\pi x)) W_{eq}(x) & \text{for } x \in (0,1), \\ W(t,0) = u_0 & \text{for } t \in (0,T). \end{cases}$

We expect:

- both schemes to converge (in time) towards the original, unperturbed steady solution;
- the DG scheme with prior to provide a **better approximation of the unperturbed steady solution** than the classical DG scheme.

PINNs as a DG prior: perturbed steady solution



PINNs as a DG prior: perturbed steady solution



PINNs as a DG prior: perturbed steady solution



We use the DG scheme to solve the advection of a Gaussian bump:

$$\begin{cases} \partial_t W + \partial_x W = aW + bW^2 & \text{for } x \in (0,1), \ t \in (0,T), \\ W(0,x) = 0.1(1 + e^{-100(x - 0.5)^2}) & \text{for } x \in (0,1), \\ W(t,0) = 0.1(1 + e^{-25}) & \text{for } t \in (0,T). \end{cases}$$

We expect the prior not to alter the convergence:

- both schemes to converge with the same error rate;
- the DG scheme with prior to provide a **similar approximation** to the classical DG scheme.

We compute the errors in x between the exact and approximate solutions:

- for several numbers of basis elements and discretization cells,
- using a = 0.75; b = 0.75; $u_0 = 0.15$.

	without	without prior		with prior			
cells	error	order		error	order	gain	
10	4.04e-02	_		5.04e-02	_	0.80	
20	3.46e-02	0.22		4.28e-02	0.24	0.81	
40	2.84e-02	0.28		3.50e-02	0.29	0.81	
80	2.15e-02	0.40		2.64e-02	0.40	0.81	
160	1.47e-02	0.55		1.81e-02	0.55	0.81	

(i) Errors with a basis composed of one element.

We compute the errors in x between the exact and approximate solutions:

- for several numbers of basis elements and discretization cells,
- using a = 0.75; b = 0.75; $u_0 = 0.15$.

	without	without prior		with prior			
cells	error	order		error	order	gain	
10	1.92e-02	_		1.93e-02	_	1.00	
20	6.26e-03	1.62		6.27e-03	1.62	1.00	
40	1.19e-03	2.39		1.20e-03	2.39	1.00	
80	1.99e-04	2.59		1.99e-04	2.59	1.00	
160	4.19e-05	2.24		4.20e-05	2.24	1.00	

(j) Errors with a basis composed of two elements.

We compute the errors in x between the exact and approximate solutions:

- for several numbers of basis elements and discretization cells,
- using a = 0.75; b = 0.75; $u_0 = 0.15$.

	without	without prior		with prior		
cells	error	order		error	order	gain
10	5.15e-03	_		5.15e-03	_	1.00
20	4.56e-04	3.50		4.56e-04	3.50	1.00
40	4.55e-05	3.32		4.55e-05	3.32	1.00
80	5.42e-06	3.07		5.42e-06	3.07	1.00
160	6.75e-07	3.01		6.75e-07	3.01	1.00

(k) Errors with a basis composed of three elements.

We compute the errors in x between the exact and approximate solutions:

- for several numbers of basis elements and discretization cells,
- using a = 0.75; b = 0.75; $u_0 = 0.15$.

	without	without prior		with prior			
cells	error	order		error	order	gain	
10	4.72e-04	_		4.72e-04	_	1.00	
20	2.87e-05	4.04		2.87e-05	4.04	1.00	
40	1.81e-06	3.99		1.81e-06	3.99	1.00	
80	1.14e-07	3.98		1.14e-07	3.98	1.00	
160	7.20e-09	3.99		7.20e-09	3.99	1.00	

(l) Errors with a basis composed of four elements.