## 1D Fast oscillations (Numerical)

$$C(t) = ar{C} + A \sin\left(rac{2\pi t}{T}
ight)$$

$$T \ll au_{linear} \sim ar{C}^{-1}$$

where we do not expect different dynamics if  $A>\bar{C}$  or  $A<\bar{C}$ .

### Kink dynamics

$$\dot{d}(t) \simeq -24\sqrt{2}ar{C}^{rac{1}{2}}e^{-2^{rac{1}{2}}ar{C}^{rac{1}{2}}d}$$

**Notice**: It is the same formula in 1D Constant (Numerical) but now  $C \to \bar{C}$ .

The variation of the distance over a period (assuming the distant to be constant inside the integrand)

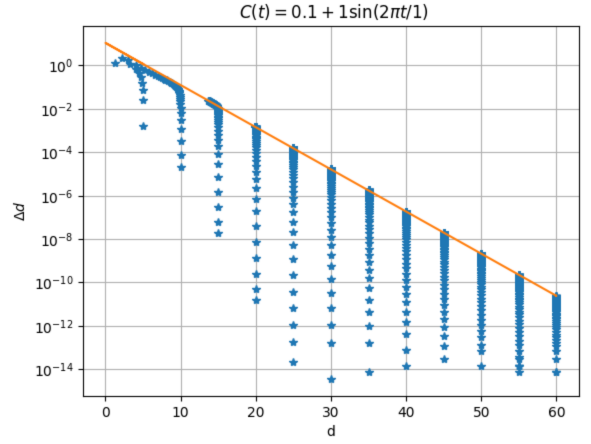
$$\Delta d(d) = \int_0^T (\partial_t d) dt = -24\sqrt{2}\int_0^T e^{d\sqrt{2}ar{C}^{1/2}}ar{C}^{1/2}dt$$

#### **Simulation**

In all the following simulations, we adopted  $T=10\bar{C}^{-1}$ , such that  $T\gg C^{-1}$ .

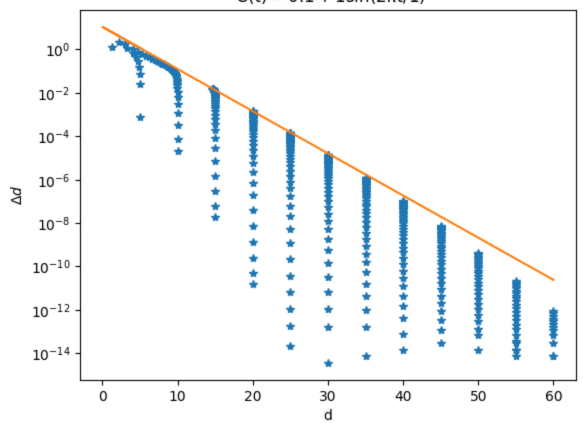
The orange line is the formula above, while the blue points are experimental values. Those values are obtained running a simulation of 100T "seconds" for many different values of the initial distance.  $\Delta d$  is calculated for each period, for each simulation.

# Variation of the distance of twokinks over one period T tspan=100\*T; dt = T/100

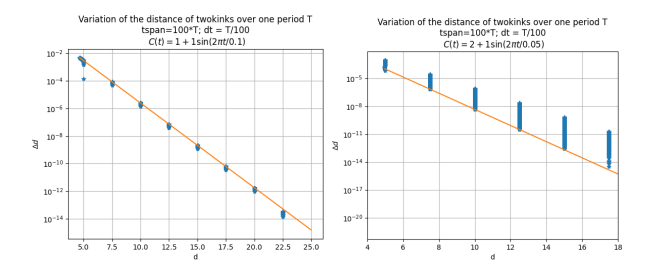


**Notice**: The way the blue dots "reach" the model tells something: We need to wait some periods before the simulation matches the model. Probably this is related to the **initial state preparation**. Here we show the same plot above, but each simulation lasts 50T instead of

# Variation of the distance of twokinks over one period T tspan=50\*T; dt = T/100 $C(t) = 0.1 + 1\sin(2\pi t/1)$



### Other simulations



## **Linear dynamics**

$$\ell = rac{2\pi}{< q^2 >^{1/2}} \sim t^{1/2}$$

# Starting from random initial state $\ell = 2\pi/ < q^2 > ^{1/2}$

