

exploit-capillary_discharge-no_bg_remove

May 22, 2025

1 Exploitation de l'expérience du plasma pulsé

```
[1]: # %matplotlib ipympl # for interactive plots

import imageio.v2 as imageio
import matplotlib.pyplot as plt
import numpy as np
import scipy.stats as stats
import scipy.integrate as si
from scipy import constants as cons
import pandas as pd
import os

import sys
sys.path.append("../..xspectra/src")
from xspectra import simulation as xs_sim
from xspectra import utils as xs_utils
from xspectra import visualization as xs_vis

from mpl_toolkits.mplot3d import Axes3D
from scipy.signal import find_peaks
from copy import deepcopy
```

1.1 Protocole

Dans l'expérience précédente, nous avons observé un plasma en émission continue. Cependant, bien des plasmas n'émettent que sur un court instant, d'où l'intérêt de cette deuxième manipulation, qui vise à observer un plasma pulsé de diazote.

Pour cela on réalise le montage suivant :

La décharge a lieu dans la cellule ci-dessous :

Ici un gaz composé à **95%** de N₂ et à **5%** de O₂ est soumis à une tension de l'ordre de **10 kV** pour générer un plasma dans une cellule de décharge. Le tout est mis dans une cage de Faraday (cf. Figure 1) afin d'atténuer les bruits électromagnétiques dans le laboratoire. À l'aide d'une fibre optique, le plasma est observé au spectroscope, au niveau de la cathode (*high*), de l'anode (*ground*) et entre les deux (*center*). Celui-ci est centré sur la raie caractéristique **C3Π(v' = 0) → B3Πg(v'' = 0)** du diazote, se trouvant à **λ = 337 nm**. Une des difficultés de l'expérience est

de déclencher la caméra au bon moment. Pour cela, on récupère d'abord les différents signaux sur l'oscilloscope pour repérer les impulsions du plasma, puis on règle manuellement le déclenchement de la caméra en ajoutant un délai à partir du déclenchement externe récupéré sur le générateur fournissant la décharge dans la cellule. Pour obtenir un meilleur signal, on augmente le nombre d'acquisitions en effectuant des décharges de manière périodique à une fréquence de **10 Hz**.

D'autre part, la pression dans l'enceinte du plasma est contrôlée à la main (faute de mieux) pour la garder autour de **27 mbar** (si la pression est trop basse, il n'y a pas d'émission, tandis que si elle est trop haute, il faut plus d'énergie pour ioniser le tout). Le débit de gaz, quant à lui, est maintenu à **50 cm³/min** à l'aide d'un contrôleur de débit fonctionnant avec une boucle de rétroaction¹. (Voir la Figure 2 pour le schéma de l'expérience).

¹ Dans le cadre de notre expérience, nous avons utilisé un produit de [Brooks Instrument](#).

1.2 Study of the tension inside the cable of alimentation

1.2.1 Load the data of the tension in the cable of alimentation

```
[2]: ## Load oscilloscope data
data = pd.read_csv('../data/2025-03-28-capillary-discharge/oscilloscope.txt',
    ↪ skiprows=4, sep='\t')
```

```
[3]: data["Ampl"]*=9.5e3/data["Ampl"].max() # Correcting the value due to the gain
    ↪ added on the experiment
data.head()
```

```
[3]:
```

	Time	Ampl
0	-1.040082e-07	-63.972969
1	-1.038082e-07	160.820794
2	-1.036082e-07	31.986520
3	-1.034082e-07	-287.878185
4	-1.032082e-07	-71.081140

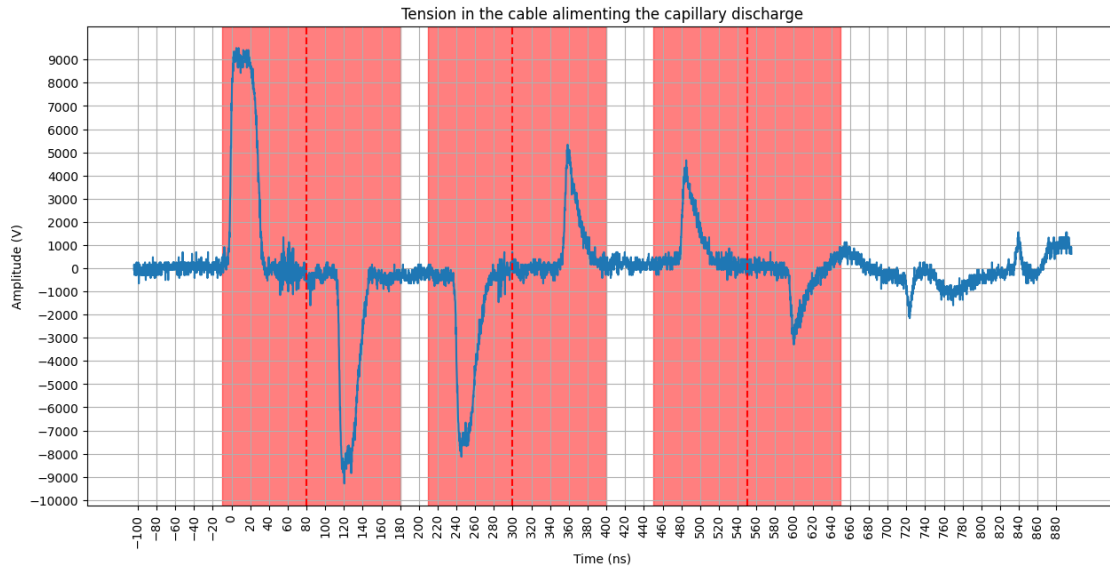
```
[4]: oscilloscopes_pulses =
    ↪ [((-10,80),(80,180)),((210,300),(300,400)),((450,550),(550,650))]
```

```
[5]: plt.figure(figsize=(15, 7))
plt.plot(data["Time"]*1e9, data["Ampl"])
plt.xlabel("Time (ns)")
plt.ylabel("Amplitude (V)")
plt.title("Tension in the cable alimenting the capillary discharge")
plt.grid()
plt.xticks(ticks=np.arange(-100,900, 20))
plt.yticks(ticks=np.arange(-10_000,10_000, 1000))
plt.xticks(rotation=90)

for pulse in oscilloscopes_pulses:
    plt.axvspan(pulse[0][0], pulse[1][1], color='red', alpha=0.5, label="Pulse")
```

```
plt.axvline(pulse[0][1], color='red', linestyle='--')

# plt.xlim((-34,100))
plt.savefig("res/oscilloscope.png")
plt.show()
```



Le signal de la tension dans le câble (cf. Figure ci-dessus) rend compte des impulsions auxquelles est soumis le plasma. Une impulsion - ou *pulse* - se réfléchit plusieurs fois sur les extrémités du câble, d'où les plusieurs échos observés, échos générant des décharges secondaires. On remarque une légère augmentation du bruit après le premier pic : ceci correspond à la première émission du plasma, d'où un délai d'une centaine de nanosecondes pour la première décharge.

1.2.2 Energie des pulses

Nous pouvons calculer les énergies dissipées dans le plasma à chaque pulse.

```
[6]: impedance_cable = 50 # Ohm
    energie_total = 0

    results = []

    for i, pulse in enumerate(oscilloscopes_pulses):
        # Filtrer les données dans l'intervalle [0, 200]
        filter_input = data[(data["Time"] * 1e9 >= pulse[0][0]) & (data["Time"] * 1e9 <= pulse[0][1])]
        filter_output = data[(data["Time"] * 1e9 >= pulse[1][0]) & (data["Time"] * 1e9 <= pulse[1][1])]
```

```

# Calculer l'intégrale en utilisant la méthode de Simpson
input = si.simpson(filter_input["Ampl"] ** 2, filter_input["Time"]) /  $\Delta$ 
↳ impedance_cable
output = si.simpson(filter_output["Ampl"] ** 2, filter_output["Time"]) /  $\Delta$ 
↳ impedance_cable
gain_energie = input - output
energie_total += gain_energie

# Ajouter les résultats au tableau
results.append({
    "Pulse": i + 1,
    "Énergie Input (mJ)": input * 1e3,
    "Énergie Output (mJ)": output * 1e3,
    "Gain Énergie (mJ)": gain_energie * 1e3
})

# Créer un DataFrame pandas
results_df = pd.DataFrame(results)

# Afficher l'énergie totale
print(f"Énergie totale : {energie_total * 1e3:4.2f} mJ")
results_df

```

Énergie totale : 35.06 mJ

```

[6]:
   Pulse  Énergie Input (mJ)  Énergie Output (mJ)  Gain Énergie (mJ)
0      1          42.851902          25.522590          17.329312
1      2          20.519548           5.118434          15.401113
2      3           4.233049           1.899990           2.333059

```

Effectuons le calcul théorique d'un échauffement homogène du gaz présent dans le tube capillaire.

```

[7]: P=27e2 # Pa
rayon_capillaire = 0.5e-3 # m
longueur_capillaire = 5e-2 # m
V=np.pi * (rayon_capillaire**2) * longueur_capillaire # m3
T1 = 300 # K
T2 = 1200 # K

n = P * V / (cons.R * T1) # mol
densite = cons.Avogadro * n / (V*10**6) # nb_particules/cm3

energie_theorique = (5/2) * n * cons.R * (T2 - T1) # J
print(f"Nombre de particules : {n:.2e} mol")
print(f"Densité : {densite:.2e} nb_particules/cm3")
print(f"Énergie théorique : {energie_theorique*1e3:4.2f} mJ")

```

Nombre de particules : 4.25×10^{-8} mol
Densité : 6.52×10^{17} nb_particules/cm³
Énergie théorique : 0.80 mJ

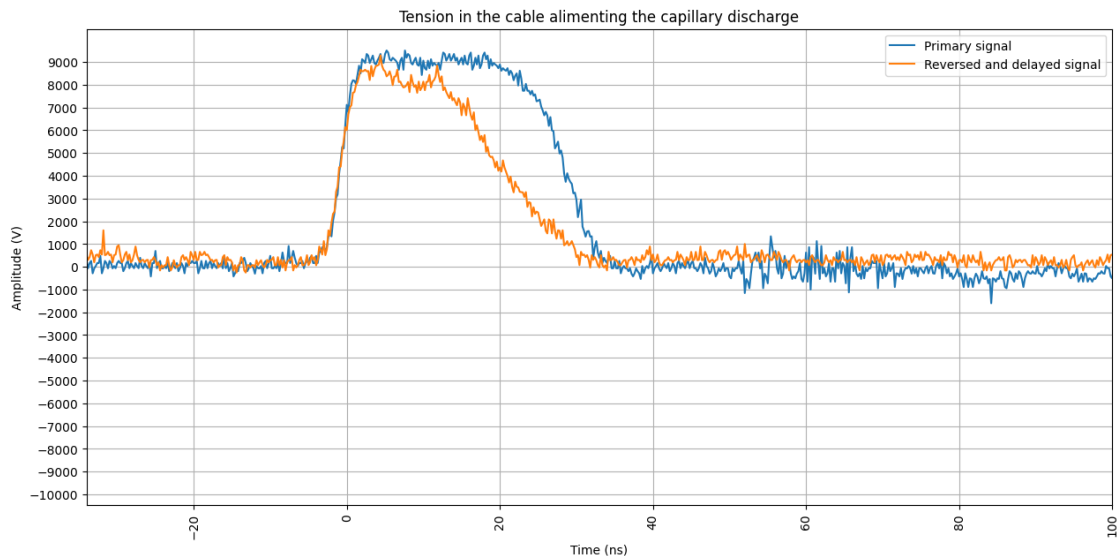
1.2.3 Instant power

```
[8]: delay=116 #ns

# Tentative d'alignement avec l'écho ....
plt.figure(figsize=(15, 7))
plt.plot(data["Time"]*1e9, data["Ampl"], label="Primary signal")
plt.plot(data["Time"]*1e9-delay, -data["Ampl"], label="Reversed and delayed ↵
↵signal")
plt.xlabel("Time (ns)")
plt.ylabel("Amplitude (V)")
plt.title("Tension in the cable alimenting the capillary discharge")
plt.grid()
plt.savefig("res/oscilloscope.png")
plt.xticks(ticks=np.arange(-100,900, 20))
plt.yticks(ticks=np.arange(-10_000,10_000, 1000))
plt.xticks(rotation=90)

plt.legend()

plt.xlim((-34,100))
plt.show()
```



```
[9]: time_step = (data["Time"].max() - data["Time"].min() ) / len(data)
delay_in_steps = int(delay * 1e-9 / time_step)

T = data["Time"].values[:-delay_in_steps]
tension = data["Ampl"].values[:-delay_in_steps]
echo = -data["Ampl"].values[delay_in_steps:]
instant_power = (tension**2 - echo**2) / impedance_cable
```

```
[10]: fig, ax1 = plt.subplots(figsize=(15, 7))

# Plot tension on the first y-axis
ax1.plot(data["Time"]*1e9, data["Ampl"], color="blue")
ax1.set_ylabel("Tension (V)", color="blue")
ax1.tick_params(axis='y', labelcolor="blue")

# Center the y-axis of ax1 at 0
max_abs_ampl = max(abs(data["Ampl"].min()), abs(data["Ampl"].max())) * 1.1
ax1.set_ylim(-max_abs_ampl, max_abs_ampl)

# Create a second y-axis for instant power
ax2 = ax1.twinx()

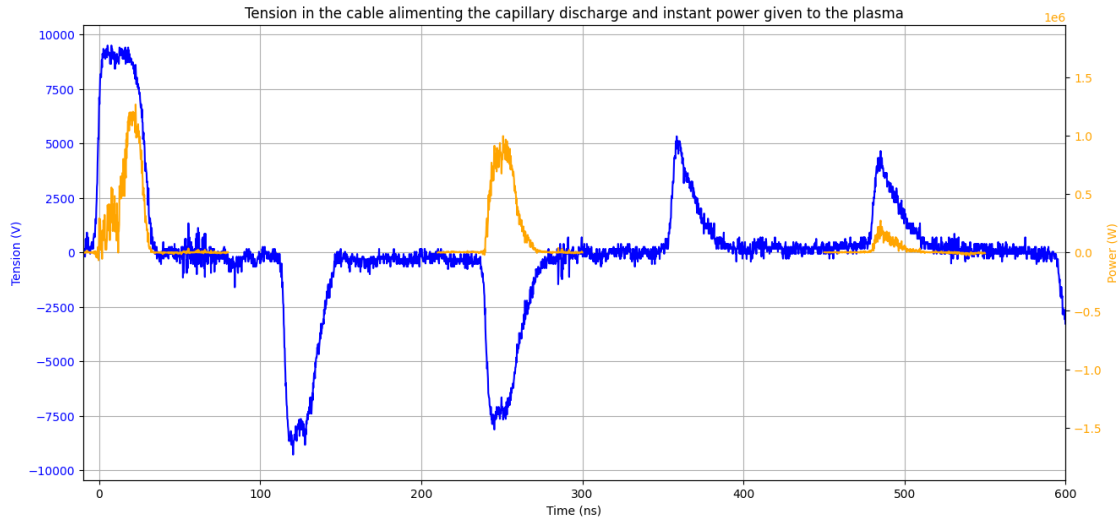
for i, pulse in enumerate(oscilloscopes_pulses):
    filter = (T*1e9 >= pulse[0][0]) & (T*1e9 <= pulse[0][1])
    ax2.plot(T[filter]*1e9, instant_power[filter], color="orange")

ax2.set_ylabel("Power (W)", color="orange")
ax2.tick_params(axis='y', labelcolor="orange")

# Center the y-axis of ax2 at 0
max_abs_power = max(abs(instant_power.min()), abs(instant_power.max())) * 1.1
ax2.set_ylim(-max_abs_power, max_abs_power)

# Add grid, title, and labels
plt.title("Tension in the cable alimentering the capillary discharge and instant_
power given to the plasma")
ax1.set_xlabel("Time (ns)")
ax1.grid()

plt.xlim((-10,600))
plt.savefig("res/instant_power.png")
plt.show()
```



```
[11]: # On clear la mémoire du jupyter pour la suite
      # %reset -f
```

1.3 Loading the atom lights used for calibration

```
[12]: # Récupération de manière séparée de la longueur d'onde
      wavelengths = xs_utils.get_winspec_spectra("../data/
      ↪2025-03-28-capillary-discharge/spectra-337.txt")["Wavelength"].to_numpy().
      ↪astype(float)
      wavelengths253 = xs_utils.get_winspec_spectra("../data/
      ↪2025-03-28-capillary-discharge/spectra-253.txt")["Wavelength"].to_numpy().
      ↪astype(float) # pour la lampe témoin
      wavelengths
```

```
[12]: array([327.39 , 327.408, 327.427, ..., 346.529, 346.548, 346.567])
```

```
[13]: # lampe_l337 = xs_utils.load_data("../data/2025-03-28-capillary-discharge/
      ↪lampes/g10_f150micro_r1200BLZ_w100ms_lampe337nm.SPE")
      lampe_l253 = xs_utils.load_data("../data/2025-03-28-capillary-discharge/lampes/
      ↪g10_f150micro_r1200BLZ_w100ms_lampe253nm.SPE")

      # Process lamp data
      # lampe_l337_filtered = xs_utils.filter_data(lampe_l337)
      lampe_l253_filtered = xs_utils.filter_data(lampe_l253,15)

      # Calculate spectra
      # spectra_337 = xs_utils.compute_spectra(lampe_l337)
      spectra_253 = xs_utils.compute_spectra(lampe_l253)
```

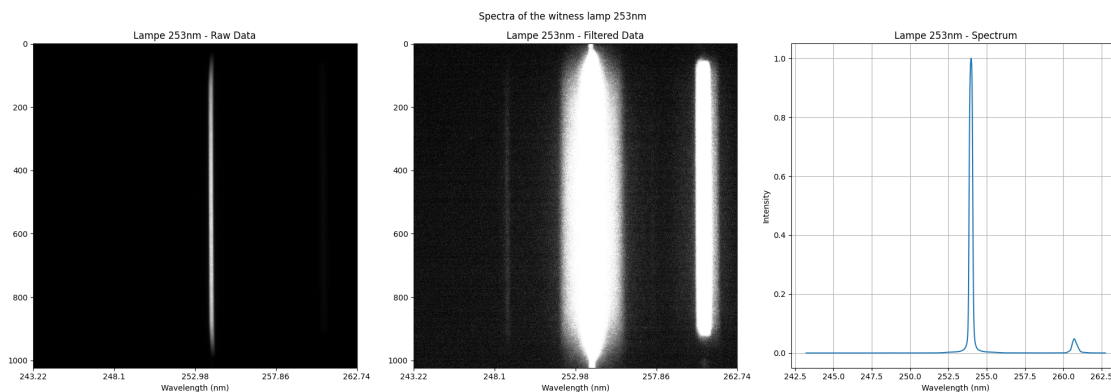
```
[14]: # Create plot
fig, axes = plt.subplots(1, 3, figsize=(20, 7))

# Plot the 253nm lamp data
axes[0].imshow(lampe_1253, cmap='gray')
axes[0].set_title("Lampe 253nm - Raw Data")
axes[0].set_xticks(ticks=np.linspace(0, lampe_1253.shape[1], num=5))
axes[0].set_xticklabels(np.linspace(wavelengths253[0], wavelengths253[-1],
    ↪ num=5).round(2))
axes[0].set_xlabel("Wavelength (nm)")

# Plot the filtered 253nm lamp data
axes[1].imshow(lampe_1253_filtered, cmap='gray')
axes[1].set_title("Lampe 253nm - Filtered Data")
axes[1].set_xticks(ticks=np.linspace(0, lampe_1253_filtered.shape[1], num=5))
axes[1].set_xticklabels(np.linspace(wavelengths253[0], wavelengths253[-1],
    ↪ num=5).round(2))
axes[1].set_xlabel("Wavelength (nm)")

# Plot the 253nm lamp spectrum
axes[2].plot(wavelengths253, spectra_253)
axes[2].set_title("Lampe 253nm - Spectrum")
axes[2].set_xlabel("Wavelength (nm)")
axes[2].set_ylabel("Intensity")
axes[2].grid(True)

plt.suptitle("Spectra of the witness lamp 253nm")
plt.tight_layout()
# plt.savefig("res/lampe_253nm.png")
plt.show()
```



On another jupyter we fit the primary ray with a gaussian of width 0.1 nm

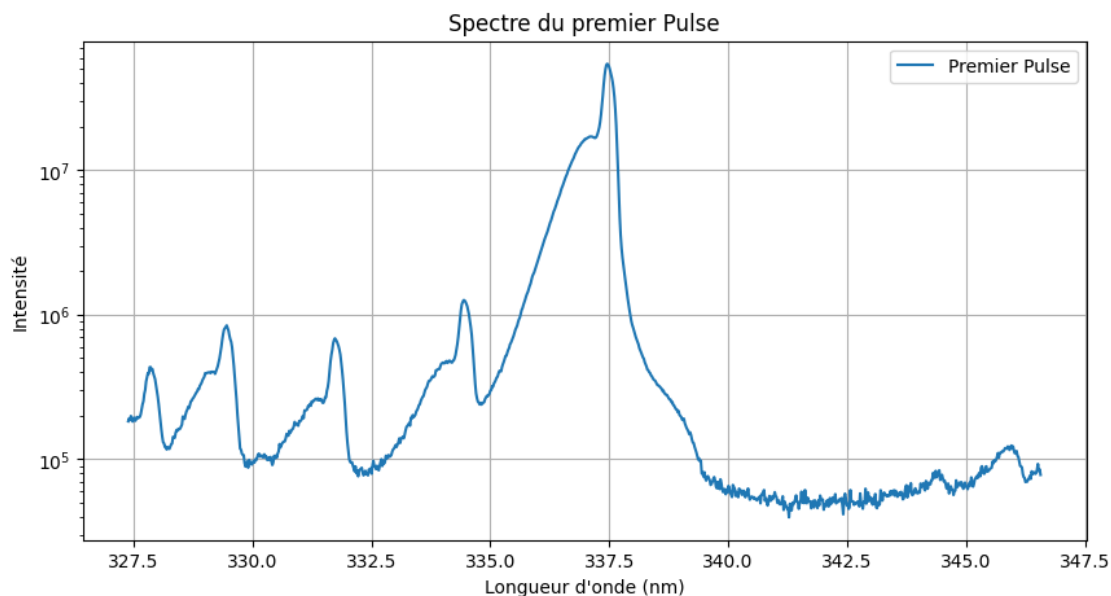
1.4 Loading of the data of the spectrum

```
[15]: center_data = xs_utils.get_points_folder("../data/
      ↪2025-03-28-capillary-discharge/center")
      ground_data = xs_utils.get_points_folder("../data/
      ↪2025-03-28-capillary-discharge/ground")
      high_data = xs_utils.get_points_folder("../data/2025-03-28-capillary-discharge/
      ↪high")
      # kyrill_data = xs_utils.get_points_folder("../data/
      ↪2025-03-28-capillary-discharge/kyrill", kyrill=True) # Data from Kyrill made
      ↪on week later
```

1.4.1 Background suppression

```
[16]: spectre_pulse_1 = xs_utils.compute_spectra(center_data[1][1], False)

plt.figure(figsize=(10, 5))
plt.plot(wavelengths, spectre_pulse_1, label="Premier Pulse")
plt.xlabel("Longueur d'onde (nm)")
plt.ylabel("Intensité")
plt.title("Spectre du premier Pulse")
plt.yscale("log")
plt.legend()
plt.grid()
plt.show()
```



```
[17]: background = center_data[1][1][:,(340 < wavelengths) & (wavelengths < 342)].
      ↪flatten()
      background.shape
```

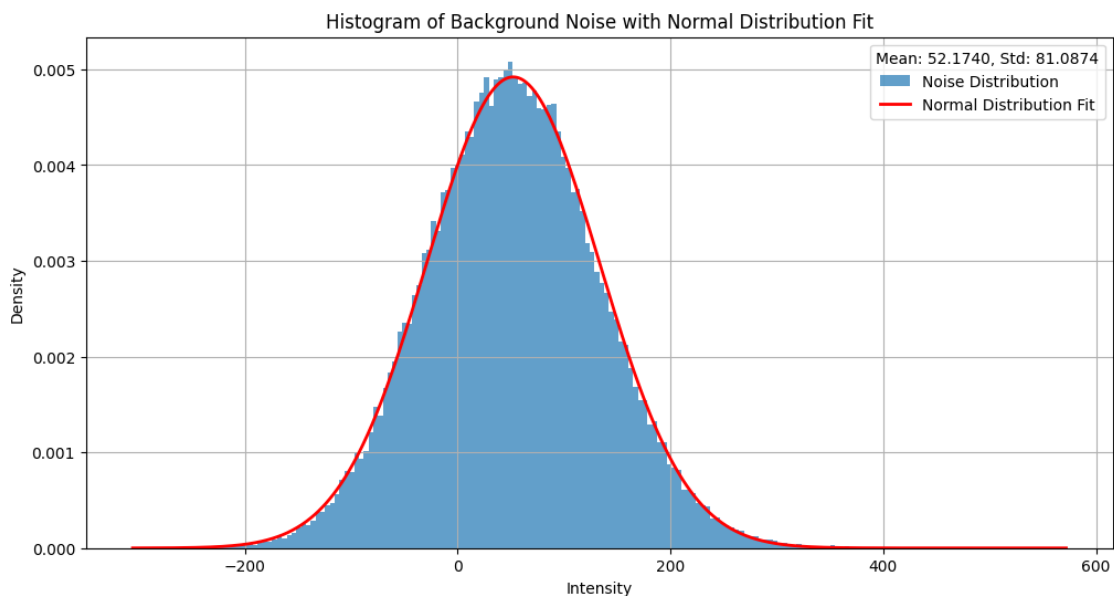
```
[17]: (109568,)
```

```
[18]: ## on affiche alors la distribution
      # Plot the histogram and fit a normal distribution
      plt.figure(figsize=(12, 6))

      # Histogram
      hist, bins, _ = plt.hist(background, bins="auto", alpha=0.7, density=True,
      ↪label='Noise Distribution')

      # Fit normal distribution
      mean = np.mean(background)
      std = np.std(background)
      x = np.linspace(min(background), max(background), 1000)
      normal_dist = stats.norm.pdf(x, mean, std)

      # Affichage de la courbe de la distribution normale
      plt.plot(x, normal_dist, 'r-', label='Normal Distribution Fit', linewidth=2)
      plt.xlabel('Intensity')
      plt.ylabel('Density')
      plt.title('Histogram of Background Noise with Normal Distribution Fit')
      plt.legend()
      plt.legend(title=f"Mean: {mean:.4f}, Std: {std:.4f}")
      plt.grid()
```



```
[19]: def remove_background(data, bgboundaries):
      """
      Remove the background from the data using the mean of the background
      """
      # Calculate the mean of the background
      mean_background = np.mean(data[:, bgboundaries[0]:bgboundaries[1]])

      # Subtract the mean background from the data
      data_corrected = data - mean_background

      return data_corrected

[20]: # plasmas_datas = {
#     "center" : (center_data[0], [remove_background(d, (340, 342)) for d in
# ↪center_data[1]]), # couple (list_delays, images)
#     "ground" : (ground_data[0], [remove_background(d, (340, 342)) for d in
# ↪ground_data[1]]),
#     "high" : (high_data[0], [remove_background(d, (340, 342)) for d in
# ↪high_data[1]])
#     # "kyrill" : (kyrill_data[0], [remove_background(d, (340, 342)) for d in
# ↪kyrill_data[1]])
# }

plasmas_datas = {
    "center" : center_data,
    "ground" : ground_data,
    "high" : high_data,
    # "kyrill" : kyrill_data
}
```

1.4.2 Correction de la calibration

Enfin, il faut veiller à corriger la calibration en transformant **wavelengths** par une transformation linéaire. Les paramètres de cette transformation peuvent être obtenus par comparaison des têtes de bandes moléculaires puis par ajustement du spectre complet - cf jupyter précédent.

```
[21]: x1, x2 = 329.5, 337.5 # tête de bande moléculaire 3->3 et 0->0 mesuré
      y1, y2 = 328.63, 337.16 # pic 3->3 et 0->0 théorique

      scale_assumption = (y2 - y1) / (x2 - x1)
      decalage_assumption = y1 - scale_assumption * x1

      print(f"Scale assumption: {scale_assumption}")
      print(f"Decalage assumption: {decalage_assumption}")
```

```
final_stretch_factor = scale_assumption
final_bias = -22.68
```

Scale assumption: 1.0662500000000037
 Decalage assumption: -22.699375000001226

```
[22]: W = final_stretch_factor * wavelengths + final_bias # wavelenths corrected
```

1.5 Quelques fonctions de traitement pour les prochains résultats

```
[23]: pulses = np.array([(120,150),(355,380),(600,610)])
pulses
```

```
[23]: array([[120, 150],
           [355, 380],
           [600, 610]])
```

```
[24]: index_pulses_center = [np.abs(plasmas_datas["center"][0]-delay).argmin() for
    ↪ delay in [130, 370, 600]]
c_image_pulses = [plasmas_datas["center"][1][i] for i in index_pulses_center]
c_spectrum_pulses = [xs_utils.compute_spectra(plasmas_datas["center"][1][i],
    ↪ True) for i in index_pulses_center]
```

```
[25]: def get_filter(data, index_pulse):
    """
    Get the filter for the data
    """
    return (pulses[index_pulse][0] <= data[0]) & (data[0] <=
    ↪ pulses[index_pulse][1])
```

```
[26]: def is_inside_pulse(delay, pulse):
    """
    Check if the data is inside the pulse
    """
    return (pulse[0] <= delay) & (delay <= pulse[1])

def is_inside_one_of_the_pulses(delay):
    """
    Check if the data is inside one of the pulses
    """
    return np.any([is_inside_pulse(delay, pulse) for pulse in pulses])
```

```
[27]: # On va faire beaucoup de graphiques pour différentes situations donc on met ça
    ↪ sous forme de fonction
def plot_spectra_and_simulation(simulation_spectra, ax_lin, ax_log,
    ↪ epsilon=1e-3, fit_areas=[], colors = ['blue', 'orange', 'green'],
    ↪ simulation_labels=None, title="Spectres et simulations", xlims=(333, 339),
    ↪ decalages=None):
```

```

if decalages is None:
    decalages = [0]*len(simulation_spectra)

# Graphique avec échelle linéaire
for i, (observed, simulated) in enumerate(zip(c_spectrum_pulses,
simulation_spectra)):
    ax_lin.plot(W, observed, label=f'Spectre {i + 1}', c=colors[i])
    if simulation_labels is not None:
        label_sim = simulation_labels[i]
        ax_lin.plot(W-decalages[i], simulated, label=label_sim, linestyle='--',
c=colors[i])

ax_lin.set_ylabel('Intensité (linéaire)')
ax_lin.set_title(title+ " échelle linéaire")
ax_lin.legend()
ax_lin.set_xlim(xlims)
ax_lin.grid()

# Graphique avec échelle logarithmique
for i, (observed, simulated) in enumerate(zip(c_spectrum_pulses,
simulation_spectra)):
    ax_log.plot(W, [x if x > epsilon else epsilon for x in observed],
label=f'Spectre {i + 1}', c=colors[i])
    if simulation_labels is not None:
        label_sim = simulation_labels[i]
        ax_log.plot(W-decalages[i], [x if x > epsilon else epsilon for x in
simulated], label=label_sim, linestyle='--', c=colors[i])

if len(fit_areas) > 0:
    for ax in [ax_lin, ax_log]:
        for i, (start, end) in enumerate(fit_areas):
            if i==0:
                ax.axvspan(start, end, color='r', alpha=0.5, label='Limites
fit')
            else:
                ax.axvspan(start, end, color='r', alpha=0.5)

ax_log.set_yscale('log')
ax_log.set_xlabel('Longueur d\onde (nm)')
ax_log.set_ylabel('Intensité (logarithmique)')
ax_log.set_title(title+" échelle logarithmique")
ax_log.legend()
ax_log.set_xlim(xlims)
ax_log.grid()

```

```
[28]: colors_places = {
        "center": "green",
        "ground": "red",
        "high": "black",
        "kyrill": "blue"
    }
```

1.6 Study of the maxima as a function of time

```
[29]: # Le câble faisant environ 25 m, on peut estimer le délai entre deux pulses

epsilon_pol = 2.25 # epsilon du polyéthylène
v = cons.c / np.sqrt(1*2.25)
L=25
D_theorique = 2*L / v * 10**9
print(f"Délai théorique entre les deux pulses: {D_theorique:.2f} ns")
```

Délai théorique entre les deux pulses: 250.17 ns

```
[30]: plt.figure(figsize=(15,7))

maxima = {}
for name_serie, (delays, data) in plasmas_datas.items():
    maxima[name_serie] = [np.max(d) for d in data]
    maxima[name_serie] = np.array(maxima[name_serie])

plt.plot(plasmas_datas["center"][0], maxima["center"], "-.", color="green",
        label="center")

plt.plot(plasmas_datas["high"][0][plasmas_datas["high"][0]<200],
        maxima["high"][plasmas_datas["high"][0]<200], "-.", color="red",
        label="high")
plt.plot(plasmas_datas["high"][0][(200<plasmas_datas["high"][0]) &
        (plasmas_datas["high"][0]<500)],
        maxima["high"][(200<plasmas_datas["high"][0]) &
        (plasmas_datas["high"][0]<500)], "-.", color="red")
plt.plot(plasmas_datas["high"][0][500<plasmas_datas["high"][0]],
        maxima["high"][500<plasmas_datas["high"][0]], "-.", color="red")

plt.plot(plasmas_datas["ground"][0][plasmas_datas["ground"][0]<200],
        maxima["ground"][plasmas_datas["ground"][0]<200], "-.", color="black",
        label="ground")
plt.plot(plasmas_datas["ground"][0][(200<plasmas_datas["ground"][0]) &
        (plasmas_datas["ground"][0]<500)],
        maxima["ground"][(200<plasmas_datas["ground"][0]) &
        (plasmas_datas["ground"][0]<500)], "-.", color="black")
```

```

plt.plot(plasmas_datos["ground"][0][500<plasmas_datos["ground"][0]],  

↳maxima["ground"][500<plasmas_datos["ground"][0]], ".-", color="black")

plt.ylabel("Intensité maximale")
plt.xlabel("Délai (ns)")

plt.title("Intensité maximale en fonction du délai")

premier_pulse=135

plt.axvline(x=premier_pulse, color="red", linestyle=":", label="pulses  

↳théoriques")
plt.axvline(x=premier_pulse+D_theorique, color="red", linestyle=":")
plt.axvline(x=premier_pulse+2*D_theorique, color="red", linestyle=":")

# Add a 10 nanoseconds scale bar
plt.hlines(y=np.max(maxima["ground"]) * 0.9, xmin=premier_pulse+D_theorique,  

↳xmax=premier_pulse +D_theorique+ 10, color="blue", linewidth=2, label="Gate  

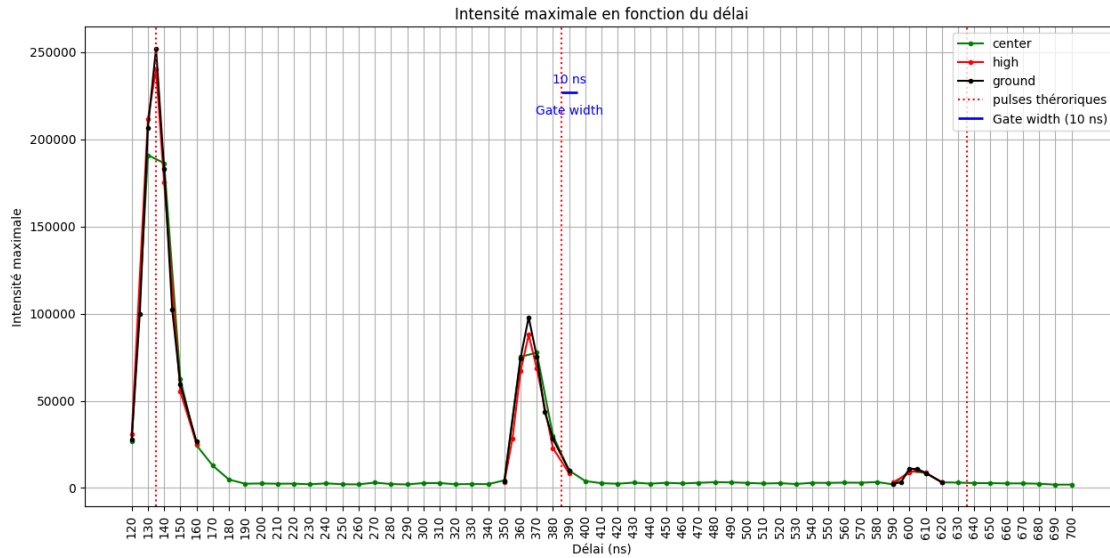
↳width (10 ns)")
plt.text(premier_pulse+D_theorique + 5, np.max(maxima["ground"]) * 0.92, "10  

↳ns", color="blue", ha="center")
plt.text(premier_pulse+D_theorique + 5, np.max(maxima["ground"]) * 0.85, "Gate  

↳width", color="blue", ha="center")

plt.grid()
plt.xticks(center_data[0], rotation=90)
plt.legend()
plt.legend()
plt.savefig("./res/intensite_max.png")
plt.show()

```



```
[31]: epsilon = 1e4 # borne inférieure pour le log
S = np.array(list(map(lambda x: xs_utils.compute_spectra(deepcopy(x), False),
    plasmas_datas["center"][1])))
S = np.array([[max(epsilon, x) for x in s] for s in S])
```

```
[32]: # Premier graphique : Échelle linéaire
fig1 = plt.figure(figsize=(10, 7))
ax1 = fig1.add_subplot(111, projection='3d')

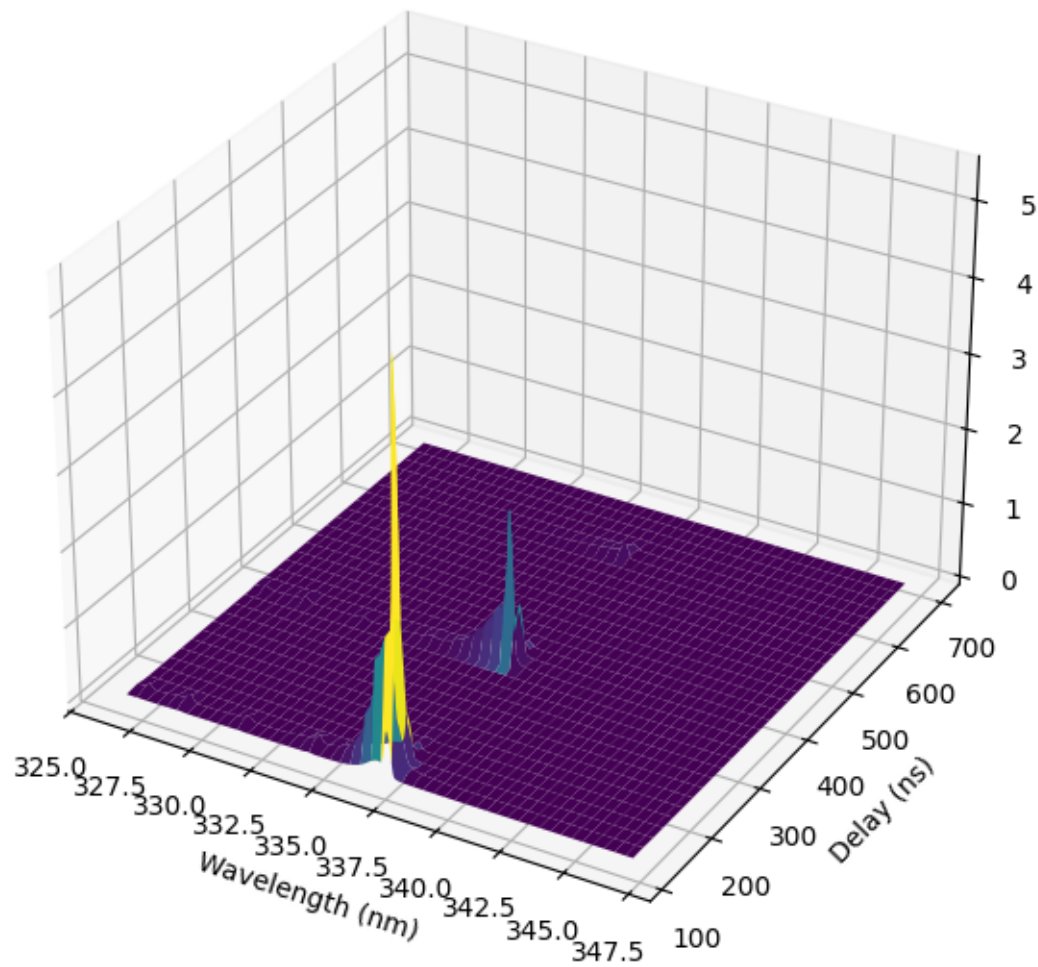
delays = plasmas_datas["center"][0]
X, Y = np.meshgrid(W, delays)
Z = S

ax1.plot_surface(X, Y, Z, cmap='viridis', edgecolor='none')
ax1.set_xlabel('Wavelength (nm)')
ax1.set_ylabel('Delay (ns)')
ax1.set_zlabel('Intensity')
ax1.set_title('Spectrum as a function of the delay - Linear Scale')

# plt.subplots_adjust(left=0.3, right=0.6)

plt.savefig("./res/3dplot_linear.png")
plt.show()
```


Spectrum as a function of the delay - Linear Scale

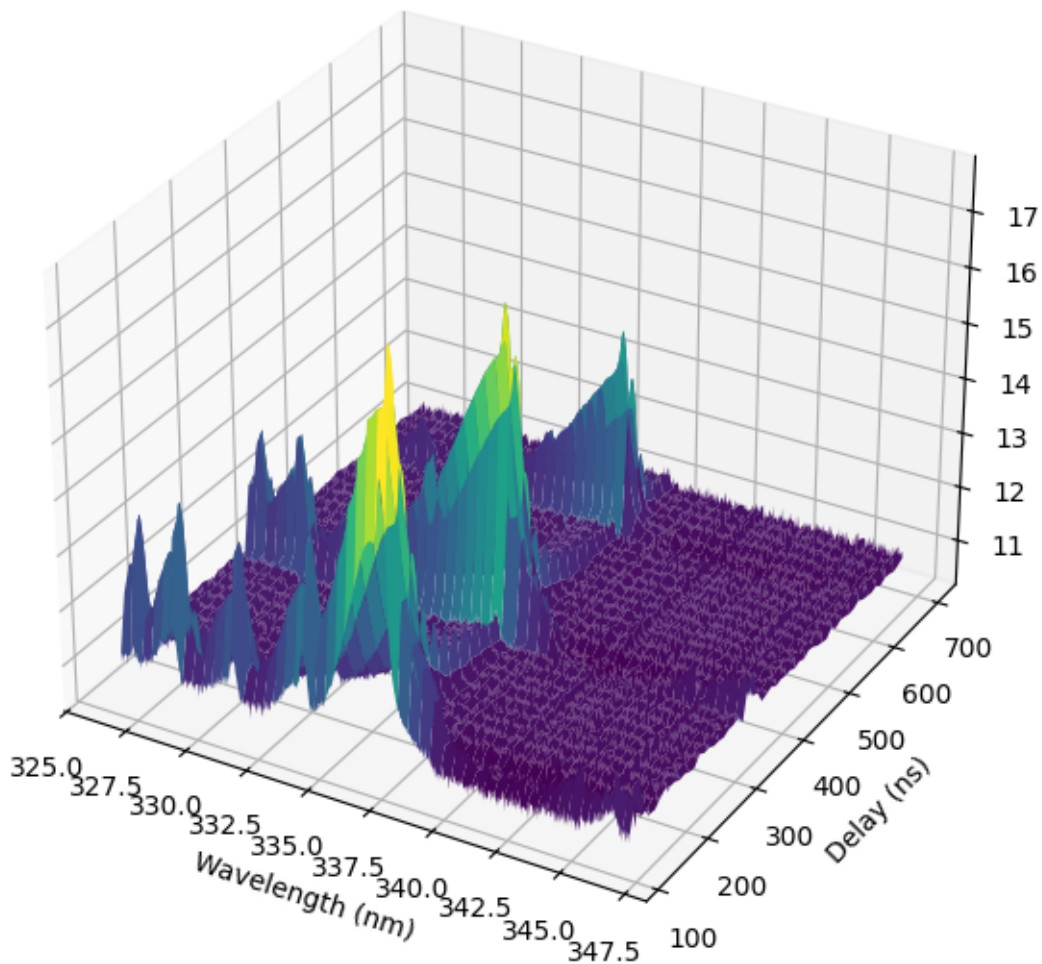


```
[33]: # Deuxième graphique : Échelle logarithmique
fig2 = plt.figure(figsize=(10, 7))
ax2 = fig2.add_subplot(111, projection='3d')

ax2.plot_surface(X, Y, np.log(Z), cmap='viridis', edgecolor='none')
ax2.set_xlabel('Wavelength (nm)')
ax2.set_ylabel('Delay (ns)')
ax2.set_zlabel('Log(Intensity)')
ax2.set_title('Spectrum as a function of the delay - Logarithmic Scale')

plt.savefig("./res/3dplot_logarithmic.png")
plt.show()
```

Spectrum as a function of the delay - Logarithmic Scale



1.7 Extraction of the temperatures

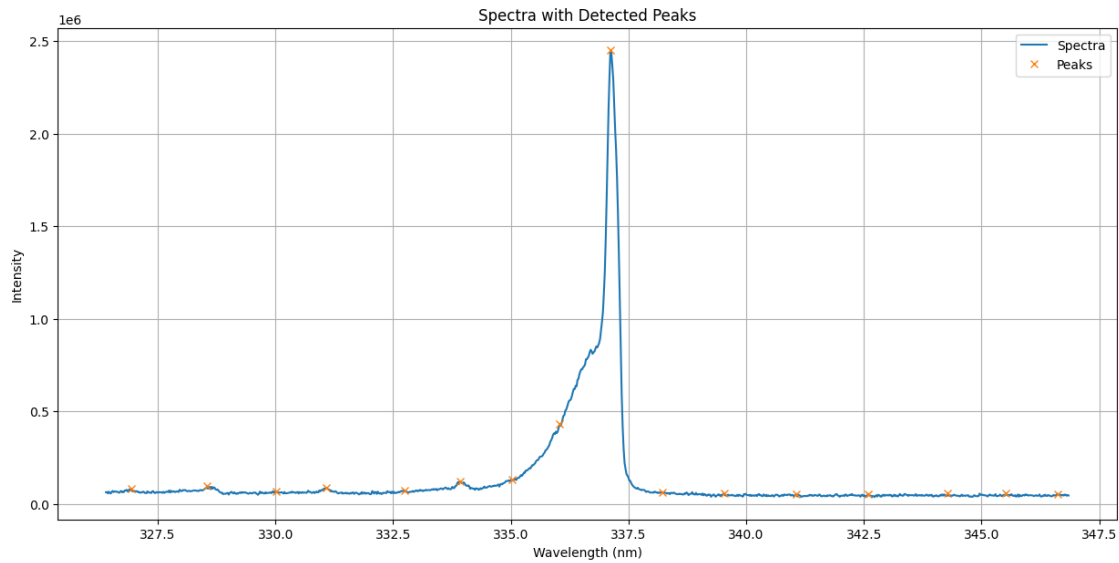
1.7.1 Température vibrationnelle par ratio

```
[34]: # Repérer les pics dans le spectre
spectra = xs_utils.compute_spectra(plasmas_datas["center"][1][5], False)
peaks, properties = find_peaks(spectra, height=0.01, distance=50)

# Afficher les résultats
plt.figure(figsize=(15, 7))
plt.plot(W, spectra, label="Spectra")
plt.plot(W[peaks], spectra[peaks], "x", label="Peaks")
plt.xlabel("Wavelength (nm)")
```

```
plt.ylabel("Intensity")
plt.title("Spectra with Detected Peaks")
plt.legend()
plt.grid()
plt.show()

# Afficher les longueurs d'onde des pics détectés
print(f"Indices des pics détectés : {peaks}")
print("Detected peaks at wavelengths:", W[peaks])
```



```
Indices des pics détectés : [ 27  107  180  233  316  375  430  481  535  590
 656  733  809  894
 956 1011]
Detected peaks at wavelengths: [326.9412425  328.547015  330.00991
 331.07296125  332.735245
 333.91665  335.01702  336.036355  337.1154  338.21470375
 339.53258875  341.069055  342.5852625  344.2784675  345.513185
 346.60822375]
```

On repère ainsi le pic dominant en 337.5nm ainsi que des pics secondaires. On prend 334.44nm comme seconde référence.

En regardant le spectre théorique, on obtient que : - le pic dominant à $\lambda_1 = 337.5\text{nm}$ correspond à la transition $C^3\Pi(\nu' = 0) \rightarrow B^3\Pi_g(\nu'' = 0)$ - le pic dominant à $\lambda_2 = 334.44\text{nm}$ correspond à la transition $C^3\Pi(\nu' = 1) \rightarrow B^3\Pi_g(\nu'' = 1)$

En suivant ainsi les formules théoriques développées dans le [jupyter de simulation](#), on a le rapport d'émission suivant :

$$r = \frac{\epsilon_1}{\epsilon_2} = \frac{n_1 \nu_1}{n_2 \nu_2} = \frac{g_{e1}(2J_1 + 1)e^{-\frac{T_{e1}}{kT_{el}} - \frac{G(\nu_1)}{kT_{vib}} - \frac{F(J_1)}{kT_{rot}}}}{g_{e2}(2J_2 + 1)e^{-\frac{T_{e2}}{kT_{el}} - \frac{G(\nu_2)}{kT_{vib}} - \frac{F(J_2)}{kT_{rot}}}} \frac{\nu_1}{\nu_2}$$

En éliminant les dégénérescences électroniques qui sont égales, ainsi que l'effet des rotations, on aboutit à :

$$r_{12} = \frac{\nu_1}{\nu_2} \exp\left(\frac{T_{e2} - T_{e1}}{kT_{el}} + \frac{G(\nu_2) - G(\nu_1)}{kT_{vib}}\right)$$

Puis, puisque l'on part du même niveau d'énergie électrique pour les deux ($C^3\Pi$), on a $T_{e1} = T_{e2}$, d'où :

$$r_{12} = \frac{\nu_1}{\nu_2} \exp\left(\frac{G(\nu_2) - G(\nu_1)}{kT_{vib}}\right)$$

Sinon on peut utiliser un troisième pic : $\lambda_3 = 331.735 \text{ nm}$ correspondant à la transition $C^3\Pi(\nu' = 2) \rightarrow B^3\Pi_g(\nu'' = 2)$ pour trouver les deux inconnues.

On a pas un spectre assez nette pour aller regarder les niveaux rotationnels.

$$T_{vib} = \frac{G(\nu_2) - G(\nu_1)}{k \ln\left(r_{12} \cdot \frac{\lambda_1}{\lambda_2}\right)}$$

En utilisant les valeurs des longueurs d'onde $\nu_1 = 337.5 \text{ nm}$ et $\nu_2 = 334.44 \text{ nm}$, ainsi que le rapport r_{12} calculé à partir des intensités des pics correspondants dans le spectre mesuré, on peut déterminer T_{vib} .

Théoriquement $G(\nu_2 = 1) - G(\nu_1 = 0) = 5.973 \times 10^{-20} - 2.019 \times 10^{-20} \text{ J} = 3.954 \times 10^{-20} \text{ J}$

Traçons l'évolution de la température.

```
[35]: i1, i2 = peaks[np.argmin(abs(W[peaks]-337.5))], peaks[np.
      ↪ argmin(abs(W[peaks]-334.4))]
```

```
i1, i2
```

```
[35]: (np.int64(535), np.int64(375))
```

```
[36]: plasmas_datas.keys()
```

```
[36]: dict_keys(['center', 'ground', 'high'])
```

```
[37]: # Calcul des températures pour chaque série
t_vib_by_ratio = {
    name : np.array([xs_utils.compute_t_vib_by_ratio(xs_utils.
      ↪ compute_spectra(d), i1=i1, i2=i2) if is_inside_one_of_the_pulses(delay) else
      ↪ 0 for delay, d in zip(delays, data)]) for name, (delays, data) in
      ↪ plasmas_datas.items()
}
```

```
# Affichage des résultats
print("Center Temperatures:", t_vib_by_ratio["center"])
```

```
Center Temperatures: [ 831.61383057  762.53265381  782.49267578  805.86065674
 0.
 0.          0.          0.          0.          0.
 0.          0.          0.          0.          0.
 0.          0.          0.          0.          0.
 0.          0.          0.          0.          885.82434082
910.20318604 952.65283203  0.          0.          0.
 0.          0.          0.          0.          0.
 0.          0.          0.          0.          0.
 0.          0.          0.          0.          0.
 0.          0.          0.          1272.7310791 1309.44042969
 0.          0.          0.          0.          0.
 0.          0.          0.          0.          ]
```

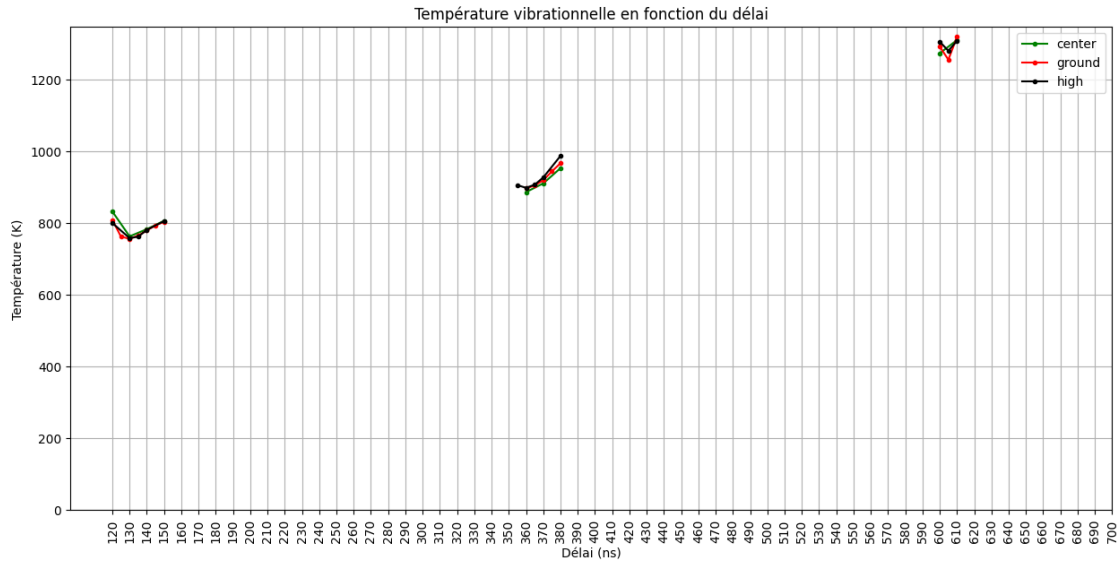
On s'aperçoit que notre méthode ne fonctionne pas du tout entre les pulses. En effet, puisqu'il n'y a pas de plasma, elle perd son sens, menant à des résultats incohérents. C'est pour cela que l'on se restreint par la suite aux pulses.

```
[38]: plt.figure(figsize=(15, 7))

# Plot center temperatures
for name in t_vib_by_ratio.keys():
    filter_1, filter_2, filter_3 = get_filter(plasmas_datas[name], 0),
    ↪ get_filter(plasmas_datas[name], 1), get_filter(plasmas_datas[name], 2)
    plt.plot(plasmas_datas[name][0][filter_1], t_vib_by_ratio[name][filter_1],
    ↪ ".-", color=colors_places[name], label=name)
    plt.plot(plasmas_datas[name][0][filter_2], t_vib_by_ratio[name][filter_2],
    ↪ ".-", color=colors_places[name])
    plt.plot(plasmas_datas[name][0][filter_3], t_vib_by_ratio[name][filter_3],
    ↪ ".-", color=colors_places[name])

plt.ylabel("Température (K)")
plt.xlabel("Délai (ns)")

plt.title("Température vibrationnelle en fonction du délai")
plt.ylim(bottom=0)
plt.grid()
plt.xticks(plasmas_datas["center"][0], rotation=90)
plt.legend()
plt.savefig("./res/temperature_vib_vs_delay.png")
plt.show()
```



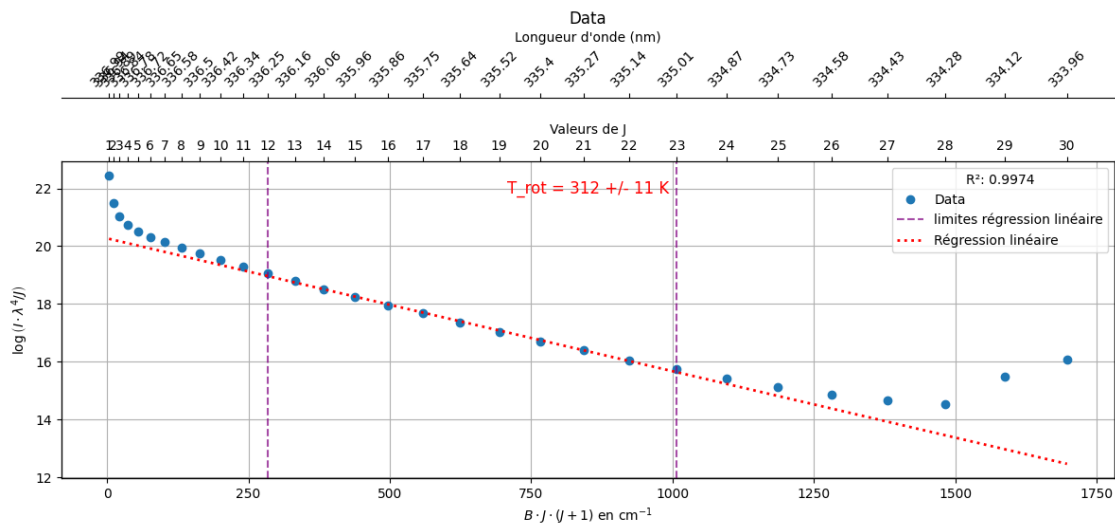
1.7.2 Extraction of the rotational temperature - Méthode analytique

```
[39]: def get_temp_rot_1(data):
    spectra = xs_utils.compute_spectra(data, True)
    return xs_utils.compute_Trot_with_branch(W, spectra, J_range=(12, 23),
    ↪certainty=0.95)

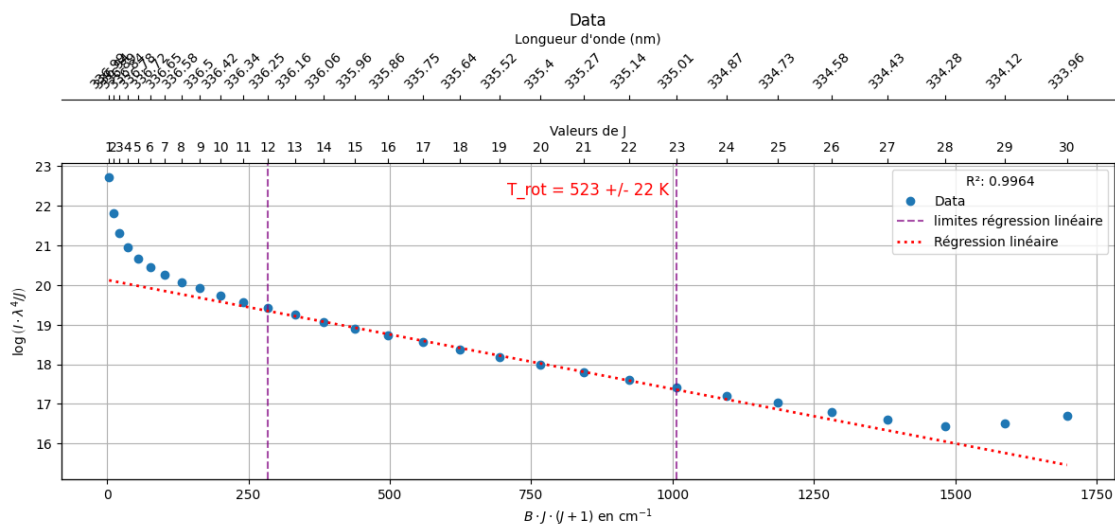
t_rot_analytic = {
    name : np.array([xs_utils.compute_Trot_with_branch(W, xs_utils.
    ↪compute_spectra(d), J_range=(12, 23), certainty=0.95) if
    ↪is_inside_one_of_the_pulses(delay) else (0,0) for delay, d in zip(delays,
    ↪data)]) for name, (delays, data) in plasmas_datas.items()
}
```

On vérifie la régression pour une valeur

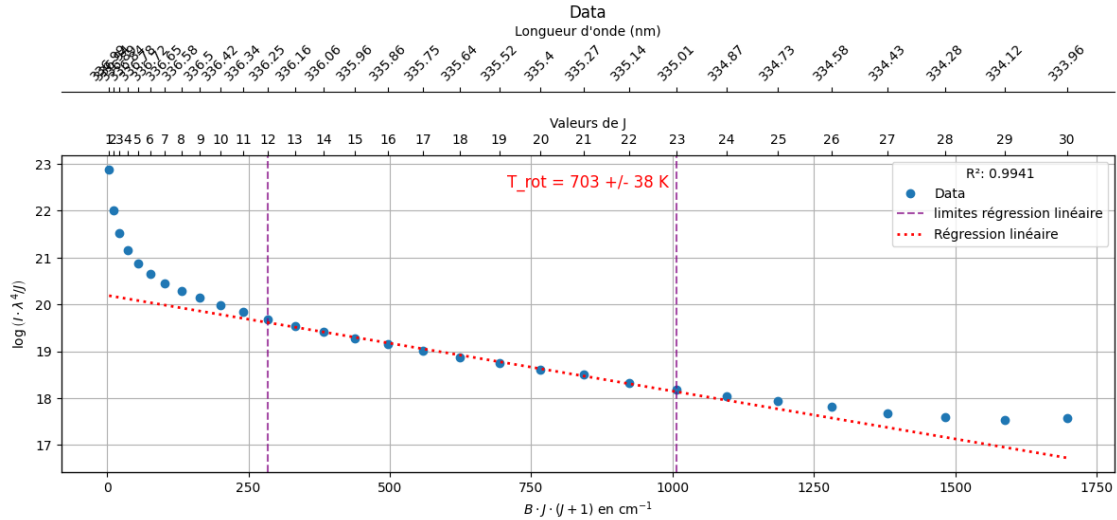
```
[40]: xs_vis.show_result_calculation_Trot(W, c_spectrum_pulses[0], J_range=(12, 23),
    ↪certainty=0.95, max_J=30) # First pulse
```



```
[41]: xs_vis.show_result_calculation_Trot(W, c_spectrum_pulses[1], J_range=(12, 23),
      ↪certainty=0.95, max_J=30) # Second pulse
```



```
[42]: xs_vis.show_result_calculation_Trot(W, c_spectrum_pulses[2], J_range=(12, 23),
      ↪certainty=0.95, max_J=30) # Third pulse
```

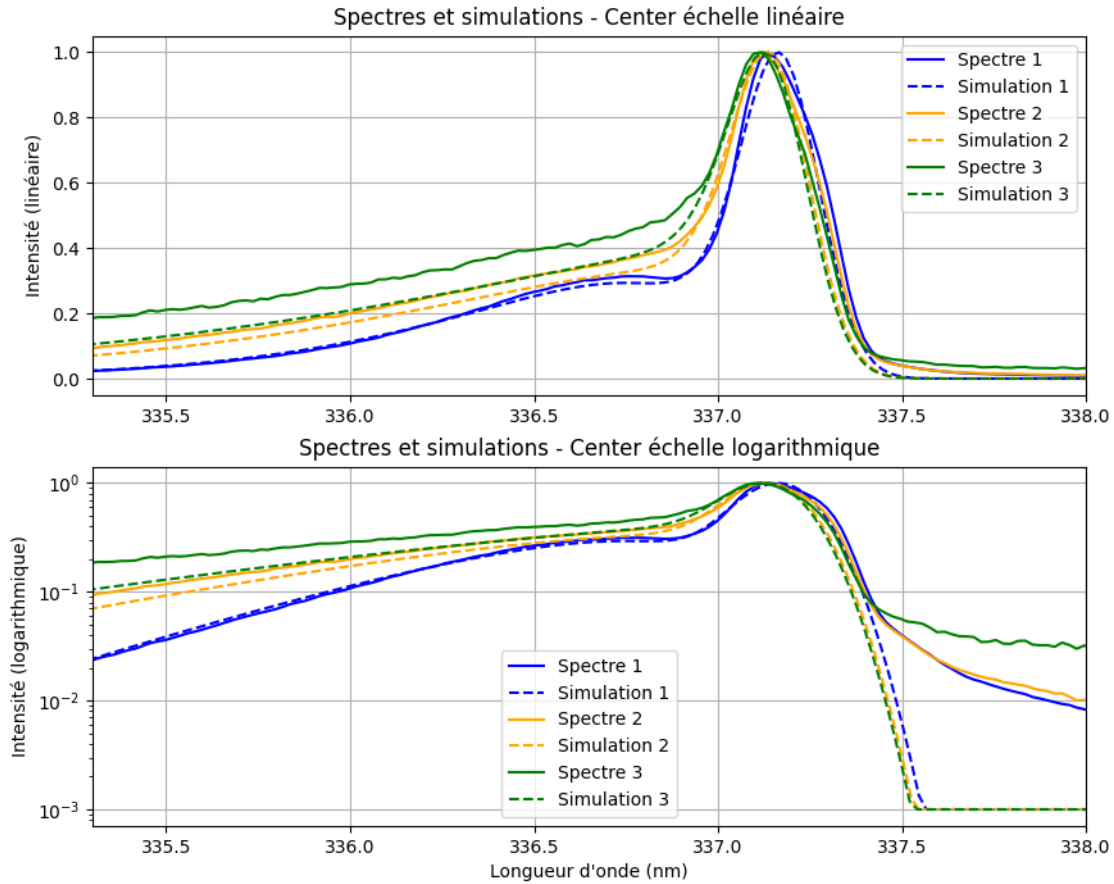


Traçons les spectres simulés à partir de ces températures pour les trois pulses du centre.

```
[43]: elargissement = 0.1
      llims = (335.3, 338)
      # Calcul du spectre de simulation
      simulation_spectra_from_reg = [xs_sim.get_spectrum(W, T_el=1_000,
      ↪T_vib=t_vib_by_ratio["center"][i], T_rot=t_rot_analytic["center"][i][0],
      ↪sigma_exp=0.1) for i in index_pulses_center]
```

```
[44]: fig, axs = plt.subplots(2, 1, figsize=(10, 8))

      plot_spectra_and_simulation(simulation_spectra_from_reg, axs[0], axs[1],
      ↪simulation_labels=["Simulation 1", "Simulation 2", "Simulation 3"],
      ↪title="Spectres et simulations - Center", decalages=[0.01,0.03,0.035],
      ↪xlims=llims)
```

Affichons les résultats en fonction du délais.

```
[45]: plt.figure(figsize=(15, 7))

for name in t_rot_analytic.keys():
    filter_1, filter_2, filter_3 = get_filter(plasmas_datas[name], 0),
    get_filter(plasmas_datas[name], 1), get_filter(plasmas_datas[name], 2)
    plt.errorbar(plasmas_datas[name][0][filter_1],
    t_rot_analytic[name][filter_1][:,0], yerr=t_rot_analytic[name][filter_1][:,1],
    fmt=".-", color=colors_places[name], label=name)
    plt.errorbar(plasmas_datas[name][0][filter_2],
    t_rot_analytic[name][filter_2][:,0], yerr=t_rot_analytic[name][filter_2][:,1],
    fmt=".-", color=colors_places[name])
    plt.errorbar(plasmas_datas[name][0][filter_3],
    t_rot_analytic[name][filter_3][:,0], yerr=t_rot_analytic[name][filter_3][:,1],
    fmt=".-", color=colors_places[name])

plt.ylabel("Température (K)")
```

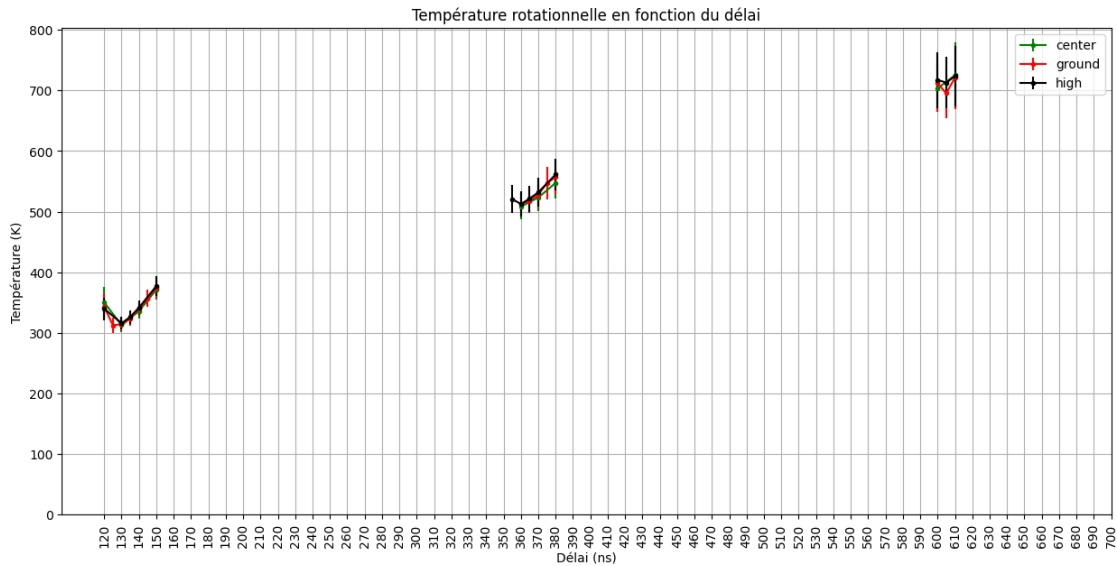
```

plt.xlabel("Délai (ns)")

plt.title("Température rotationnelle en fonction du délai")

plt.grid()
plt.xticks(plasmas_datas["center"][0], rotation=90)
plt.legend()
plt.ylim(bottom=0)
plt.savefig("./res/temperature_rot1_vs_delay.png")
plt.show()

```



1.7.3 Extraction of the rotational temperature - Fit on the $v'=0 \rightarrow v''=0$ transition

```

[46]: llims = (335, 337.5)
mask = (W > llims[0]) & (W < llims[1])

```

```

[47]: def get_temp_rot_2(data, T_vib, nb_points=None, method="sum_squares",
    ↪mask=mask):
    spectra = xs_utils.compute_spectra(data, True)
    if nb_points is not None:
        X = np.linspace(W[mask].min(), W[mask].max(), nb_points)
        Y = np.interp(X, W[mask], spectra[mask])
    else :
        X = W[mask]
        Y = spectra[mask]
    return xs_utils.get_best_fit(X, Y,
                                T_vib=T_vib,
                                elargissement=0.1,

```

```

        w_decalage=0,
        T_rot_range=(100, 2000),
        # elargissement_range=(0.05,0.12),
        w_decalage_range=(-0.5, 0.5),
        verbose=True,
        nb_steps=3,
        score_method=method)[2]

```

```

[48]: # Prends quelques minutes à exécuter
# On optimise déjà en ne prenant que les données à l'intérieur des pulses
# on pourrait également prendre moins de points # mais prend du temps même avec
↳seulement 100 points
t_rot_by_fit_simple = {
    name : np.array([get_temp_rot_2(d, t_vib_by_ratio[name][i]) if
↳is_inside_one_of_the_pulses(delay) else 0 for delay, d in zip(delays,
↳data)]) for name, (delays, data) in plasmas_datas.items()
}

```

```

Iteration  1 | Score:    0.068 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    340 K | Scale 1.00000000 | Décalage: 0.0029 nm
Iteration  2 | Score:    0.067 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    330 K | Scale 1.00000000 | Décalage: 0.0036 nm
Iteration  3 | Score:    0.067 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    327 K | Scale 1.00000000 | Décalage: 0.0039 nm
Iteration  1 | Score:    0.072 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    335 K | Scale 1.00000000 | Décalage: 0.0023 nm
Iteration  2 | Score:    0.071 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    327 K | Scale 1.00000000 | Décalage: 0.0029 nm
Iteration  3 | Score:    0.071 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    325 K | Scale 1.00000000 | Décalage: 0.0030 nm
Iteration  1 | Score:    0.073 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    378 K | Scale 1.00000000 | Décalage: 0.0024 nm
Iteration  2 | Score:    0.073 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    369 K | Scale 1.00000000 | Décalage: 0.0030 nm
Iteration  3 | Score:    0.073 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    367 K | Scale 1.00000000 | Décalage: 0.0032 nm
Iteration  1 | Score:    0.068 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    429 K | Scale 1.00000000 | Décalage: 0.0015 nm
Iteration  2 | Score:    0.067 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    423 K | Scale 1.00000000 | Décalage: 0.0018 nm
Iteration  3 | Score:    0.067 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    421 K | Scale 1.00000000 | Décalage: 0.0019 nm
Iteration  1 | Score:    0.047 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    676 K | Scale 1.00000000 | Décalage: 0.0056 nm
Iteration  2 | Score:    0.045 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    661 K | Scale 1.00000000 | Décalage: 0.0064 nm
Iteration  3 | Score:    0.045 | Elargissement:    0.10 nm | T_vib =    782 |
T_rot:    658 K | Scale 1.00000000 | Décalage: 0.0065 nm

```

Iteration 1 | Score: 0.049 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 715 K | Scale 1.00000000 | Décalage: 0.0060 nm
 Iteration 2 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 698 K | Scale 1.00000000 | Décalage: 0.0069 nm
 Iteration 3 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 696 K | Scale 1.00000000 | Décalage: 0.0070 nm
 Iteration 1 | Score: 0.049 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 755 K | Scale 1.00000000 | Décalage: 0.0062 nm
 Iteration 2 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 737 K | Scale 1.00000000 | Décalage: 0.0071 nm
 Iteration 3 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 734 K | Scale 1.00000000 | Décalage: 0.0072 nm
 Iteration 1 | Score: 0.048 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 1199 K | Scale 1.00000000 | Décalage: 0.0144 nm
 Iteration 2 | Score: 0.045 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 1164 K | Scale 1.00000000 | Décalage: 0.0154 nm
 Iteration 3 | Score: 0.045 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 1161 K | Scale 1.00000000 | Décalage: 0.0154 nm
 Iteration 1 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 1292 K | Scale 1.00000000 | Décalage: 0.0118 nm
 Iteration 2 | Score: 0.042 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 1264 K | Scale 1.00000000 | Décalage: 0.0124 nm
 Iteration 3 | Score: 0.042 | Elargissement: 0.10 nm | T_vib = 782 |
 T_rot: 1263 K | Scale 1.00000000 | Décalage: 0.0124 nm
 Iteration 1 | Score: 0.118 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 437 K | Scale 1.00000000 | Décalage: 0.0247 nm
 Iteration 2 | Score: 0.071 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 350 K | Scale 1.00000000 | Décalage: 0.0297 nm
 Iteration 3 | Score: 0.069 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 331 K | Scale 1.00000000 | Décalage: 0.0308 nm
 Iteration 1 | Score: 0.124 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 425 K | Scale 1.00000000 | Décalage: 0.0232 nm
 Iteration 2 | Score: 0.080 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 343 K | Scale 1.00000000 | Décalage: 0.0277 nm
 Iteration 3 | Score: 0.078 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 324 K | Scale 1.00000000 | Décalage: 0.0289 nm
 Iteration 1 | Score: 0.117 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 428 K | Scale 1.00000000 | Décalage: 0.0221 nm
 Iteration 2 | Score: 0.077 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 351 K | Scale 1.00000000 | Décalage: 0.0263 nm
 Iteration 3 | Score: 0.074 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 332 K | Scale 1.00000000 | Décalage: 0.0273 nm
 Iteration 1 | Score: 0.116 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 447 K | Scale 1.00000000 | Décalage: 0.0218 nm
 Iteration 2 | Score: 0.079 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 372 K | Scale 1.00000000 | Décalage: 0.0257 nm
 Iteration 3 | Score: 0.077 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 354 K | Scale 1.00000000 | Décalage: 0.0268 nm

Iteration 1 | Score: 0.115 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 473 K | Scale 1.00000000 | Décalage: 0.0218 nm
 Iteration 2 | Score: 0.080 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 399 K | Scale 1.00000000 | Décalage: 0.0258 nm
 Iteration 3 | Score: 0.078 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 382 K | Scale 1.00000000 | Décalage: 0.0267 nm
 Iteration 1 | Score: 0.113 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 500 K | Scale 1.00000000 | Décalage: 0.0232 nm
 Iteration 2 | Score: 0.076 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 420 K | Scale 1.00000000 | Décalage: 0.0275 nm
 Iteration 3 | Score: 0.075 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 407 K | Scale 1.00000000 | Décalage: 0.0283 nm
 Iteration 1 | Score: 0.105 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 517 K | Scale 1.00000000 | Décalage: 0.0232 nm
 Iteration 2 | Score: 0.070 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 439 K | Scale 1.00000000 | Décalage: 0.0274 nm
 Iteration 3 | Score: 0.069 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 427 K | Scale 1.00000000 | Décalage: 0.0281 nm
 Iteration 1 | Score: 0.072 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 734 K | Scale 1.00000000 | Décalage: 0.0243 nm
 Iteration 2 | Score: 0.049 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 665 K | Scale 1.00000000 | Décalage: 0.0276 nm
 Iteration 3 | Score: 0.048 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 656 K | Scale 1.00000000 | Décalage: 0.0281 nm
 Iteration 1 | Score: 0.069 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 750 K | Scale 1.00000000 | Décalage: 0.0240 nm
 Iteration 2 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 683 K | Scale 1.00000000 | Décalage: 0.0271 nm
 Iteration 3 | Score: 0.046 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 673 K | Scale 1.00000000 | Décalage: 0.0276 nm
 Iteration 1 | Score: 0.071 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 777 K | Scale 1.00000000 | Décalage: 0.0241 nm
 Iteration 2 | Score: 0.050 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 710 K | Scale 1.00000000 | Décalage: 0.0272 nm
 Iteration 3 | Score: 0.050 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 700 K | Scale 1.00000000 | Décalage: 0.0277 nm
 Iteration 1 | Score: 0.064 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 798 K | Scale 1.00000000 | Décalage: 0.0245 nm
 Iteration 2 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 731 K | Scale 1.00000000 | Décalage: 0.0276 nm
 Iteration 3 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 721 K | Scale 1.00000000 | Décalage: 0.0281 nm
 Iteration 1 | Score: 0.069 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 809 K | Scale 1.00000000 | Décalage: 0.0252 nm
 Iteration 2 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 739 K | Scale 1.00000000 | Décalage: 0.0285 nm
 Iteration 3 | Score: 0.046 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 729 K | Scale 1.00000000 | Décalage: 0.0290 nm

Iteration 1 | Score: 0.056 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1243 K | Scale 1.00000000 | Décalage: 0.0282 nm
 Iteration 2 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1170 K | Scale 1.00000000 | Décalage: 0.0295 nm
 Iteration 3 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1167 K | Scale 1.00000000 | Décalage: 0.0296 nm
 Iteration 1 | Score: 0.060 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1206 K | Scale 1.00000000 | Décalage: 0.0295 nm
 Iteration 2 | Score: 0.046 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1131 K | Scale 1.00000000 | Décalage: 0.0310 nm
 Iteration 3 | Score: 0.046 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1128 K | Scale 1.00000000 | Décalage: 0.0311 nm
 Iteration 1 | Score: 0.068 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1278 K | Scale 1.00000000 | Décalage: 0.0291 nm
 Iteration 2 | Score: 0.055 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1205 K | Scale 1.00000000 | Décalage: 0.0305 nm
 Iteration 3 | Score: 0.055 | Elargissement: 0.10 nm | T_vib = 756 |
 T_rot: 1202 K | Scale 1.00000000 | Décalage: 0.0306 nm
 Iteration 1 | Score: 0.119 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 440 K | Scale 1.00000000 | Décalage: 0.0247 nm
 Iteration 2 | Score: 0.072 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 354 K | Scale 1.00000000 | Décalage: 0.0296 nm
 Iteration 3 | Score: 0.070 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0307 nm
 Iteration 1 | Score: 0.123 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 440 K | Scale 1.00000000 | Décalage: 0.0228 nm
 Iteration 2 | Score: 0.081 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 360 K | Scale 1.00000000 | Décalage: 0.0272 nm
 Iteration 3 | Score: 0.079 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 341 K | Scale 1.00000000 | Décalage: 0.0283 nm
 Iteration 1 | Score: 0.119 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 455 K | Scale 1.00000000 | Décalage: 0.0226 nm
 Iteration 2 | Score: 0.080 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 378 K | Scale 1.00000000 | Décalage: 0.0268 nm
 Iteration 3 | Score: 0.078 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 360 K | Scale 1.00000000 | Décalage: 0.0278 nm
 Iteration 1 | Score: 0.122 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 486 K | Scale 1.00000000 | Décalage: 0.0229 nm
 Iteration 2 | Score: 0.084 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 405 K | Scale 1.00000000 | Décalage: 0.0273 nm
 Iteration 3 | Score: 0.083 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 391 K | Scale 1.00000000 | Décalage: 0.0281 nm
 Iteration 1 | Score: 0.105 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 525 K | Scale 1.00000000 | Décalage: 0.0239 nm
 Iteration 2 | Score: 0.069 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 446 K | Scale 1.00000000 | Décalage: 0.0281 nm
 Iteration 3 | Score: 0.068 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 433 K | Scale 1.00000000 | Décalage: 0.0289 nm

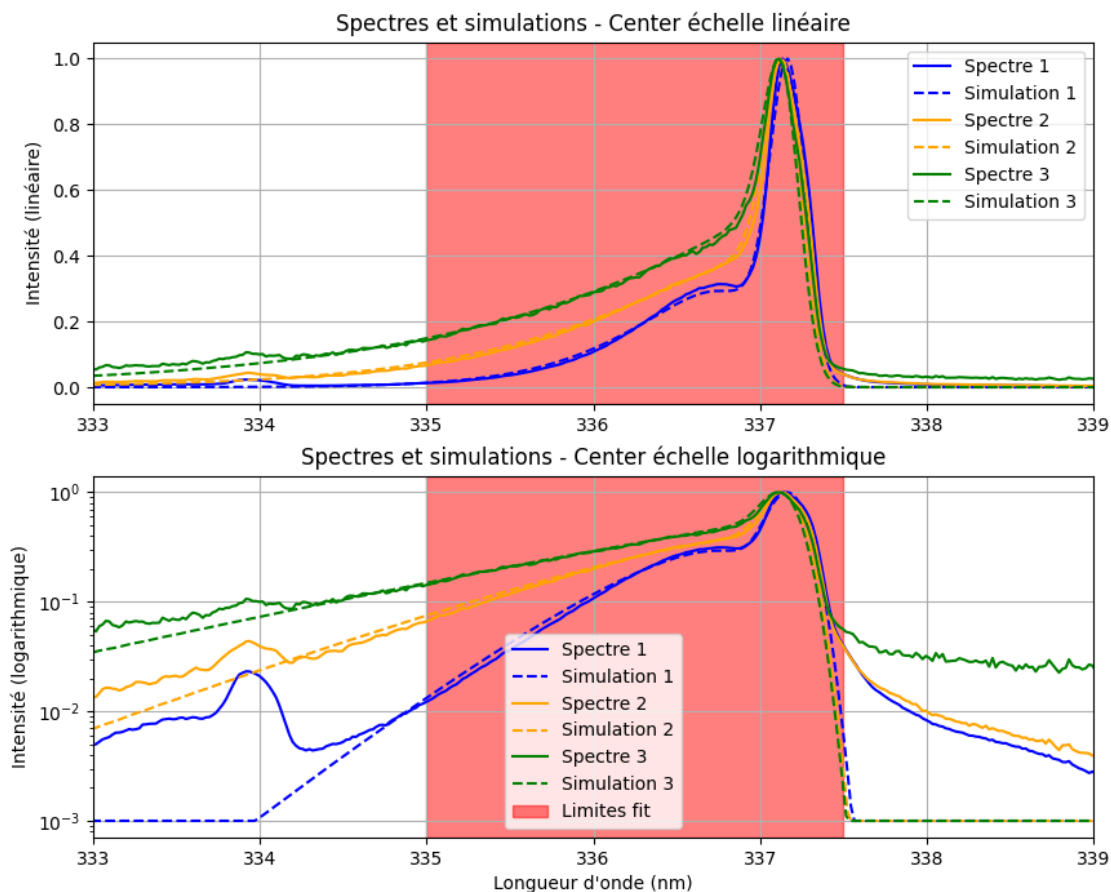
Iteration 1 | Score: 0.075 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 732 K | Scale 1.00000000 | Décalage: 0.0256 nm
 Iteration 2 | Score: 0.049 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 660 K | Scale 1.00000000 | Décalage: 0.0293 nm
 Iteration 3 | Score: 0.048 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 649 K | Scale 1.00000000 | Décalage: 0.0299 nm
 Iteration 1 | Score: 0.075 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 747 K | Scale 1.00000000 | Décalage: 0.0254 nm
 Iteration 2 | Score: 0.050 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 675 K | Scale 1.00000000 | Décalage: 0.0289 nm
 Iteration 3 | Score: 0.049 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 664 K | Scale 1.00000000 | Décalage: 0.0295 nm
 Iteration 1 | Score: 0.072 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 764 K | Scale 1.00000000 | Décalage: 0.0254 nm
 Iteration 2 | Score: 0.048 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 693 K | Scale 1.00000000 | Décalage: 0.0288 nm
 Iteration 3 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 683 K | Scale 1.00000000 | Décalage: 0.0294 nm
 Iteration 1 | Score: 0.074 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 795 K | Scale 1.00000000 | Décalage: 0.0254 nm
 Iteration 2 | Score: 0.051 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 724 K | Scale 1.00000000 | Décalage: 0.0288 nm
 Iteration 3 | Score: 0.050 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 714 K | Scale 1.00000000 | Décalage: 0.0293 nm
 Iteration 1 | Score: 0.068 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 842 K | Scale 1.00000000 | Décalage: 0.0260 nm
 Iteration 2 | Score: 0.044 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 770 K | Scale 1.00000000 | Décalage: 0.0294 nm
 Iteration 3 | Score: 0.044 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 759 K | Scale 1.00000000 | Décalage: 0.0300 nm
 Iteration 1 | Score: 0.062 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1275 K | Scale 1.00000000 | Décalage: 0.0285 nm
 Iteration 2 | Score: 0.050 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1201 K | Scale 1.00000000 | Décalage: 0.0300 nm
 Iteration 3 | Score: 0.050 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1198 K | Scale 1.00000000 | Décalage: 0.0300 nm
 Iteration 1 | Score: 0.055 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1276 K | Scale 1.00000000 | Décalage: 0.0281 nm
 Iteration 2 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1204 K | Scale 1.00000000 | Décalage: 0.0295 nm
 Iteration 3 | Score: 0.043 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1201 K | Scale 1.00000000 | Décalage: 0.0295 nm
 Iteration 1 | Score: 0.058 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1311 K | Scale 1.00000000 | Décalage: 0.0276 nm
 Iteration 2 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1240 K | Scale 1.00000000 | Décalage: 0.0288 nm
 Iteration 3 | Score: 0.047 | Elargissement: 0.10 nm | T_vib = 762 |
 T_rot: 1238 K | Scale 1.00000000 | Décalage: 0.0289 nm

Affichons les fits pour les trois pulses au centre

```
[49]: elargissement = 0.09
# Calcul du spectre de simulation
simulation_spectra_fit_sample = [xs_sim.get_spectrum(W, T_el=1_000,
    ↳T_vib=t_vib_by_ratio["center"][i], T_rot=t_rot_by_fit_simple["center"][i],
    ↳sigma_exp=0.1) for i in index_pulses_center]

[50]: fig, axs = plt.subplots(2, 1, figsize=(10, 8))

plot_spectra_and_simulation(simulation_spectra_fit_sample, axs[0], axs[1],
    ↳fit_areas=[llimits], simulation_labels=["Simulation 1", "Simulation 2",
    ↳"Simulation 3"], title="Spectres et simulations - Center", decalages=[0.01,0.
    ↳03,0.04])
```



Testons différents paramètres pour le fit pour les trois pulses.

```
[51]: plt.figure(figsize=(15, 7))

for name in t_rot_by_fit_simple.keys():
```



```

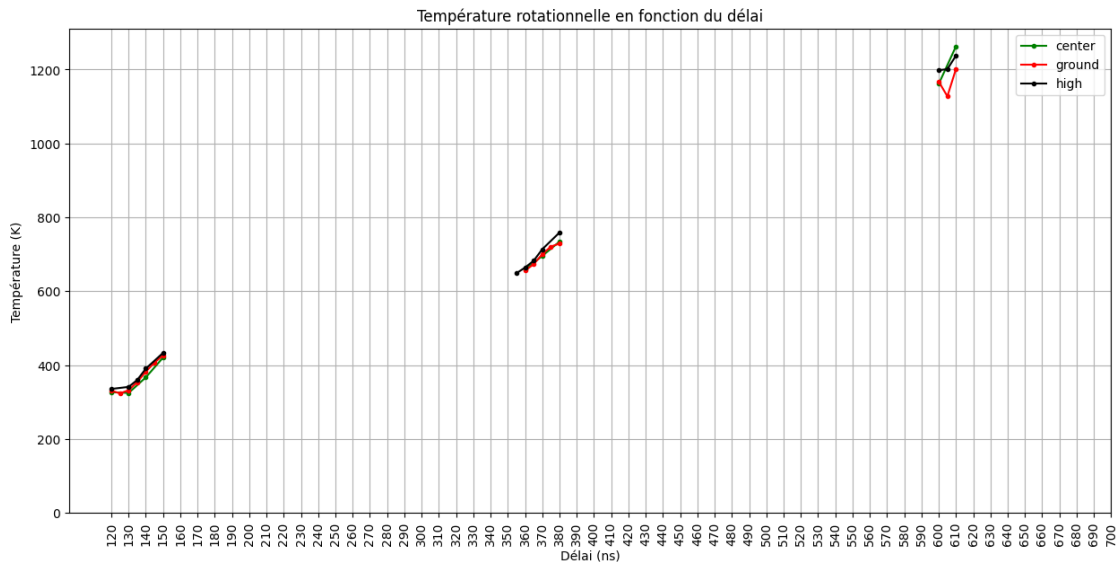
    filter_1, filter_2, filter_3 = get_filter(plasmas_datas[name], 0),
    ↪get_filter(plasmas_datas[name], 1), get_filter(plasmas_datas[name], 2)
    plt.plot(plasmas_datas[name][0][filter_1],
    ↪t_rot_by_fit_simple[name][filter_1], "-.", color=colors_places[name],
    ↪label=name)
    plt.plot(plasmas_datas[name][0][filter_2],
    ↪t_rot_by_fit_simple[name][filter_2], "-.", color=colors_places[name])
    plt.plot(plasmas_datas[name][0][filter_3],
    ↪t_rot_by_fit_simple[name][filter_3], "-.", color=colors_places[name])

plt.ylabel("Température (K)")
plt.xlabel("Délai (ns)")

plt.title("Température rotationnelle en fonction du délai")

plt.grid()
plt.xticks(plasmas_datas["center"][0], rotation=90)
plt.legend()
plt.ylim(bottom=0)
plt.savefig("./res/temperature_rot2_vs_delay.png")
plt.show()

```



1.7.4 Fit complet

```
[71]: areas = [
    (335.1, 337.6),
    (333, 334.1),
    (329.7, 331.4),
    (327.5, 328.8)
]

# areas = areas[:2]

filters_trans_vib = [(areas[i][0] <= W) & (W <= areas[i][1]) for i in
    ↪range(len(areas))]

filter_whole_fit = np.sum(filters_trans_vib, axis=0).astype(bool)

[83]: def process_whole_fit(spectrum):
    return xs_utils.get_best_fit(W[filter_whole_fit],
    ↪spectrum[filter_whole_fit],
        T_rot=400,
        T_vib=1500,
        elargissement=0.1,
        w_decalage=0, # decalage_assumption
        w_scale=1,
        T_rot_range=(300, 2_000),
        T_vib_range=(300, 10_000), # Il faut une plage
    ↪assez grande pour que la fonction soit bien convexe
        elargissement_range=(0.05,0.15),
        # w_decalage_range=(-2,2),
        # w_scale_range=(0.99,1.01), # ne pas trop
    ↪grand sinon la fonction n'est pas convexe
        modelisation_spectrum_function=xs_sim.
    ↪get_whole_spectrum, # cette fois-ci on génère les autres transitions
    ↪vibrationnelles
        verbose=True,
        nb_steps=3)

[84]: params_fit_center_pulses = np.array([process_whole_fit(c_spectrum_pulses[i])
    ↪for i in range(len(c_spectrum_pulses))])
```

```
Iteration 1 | Score: 0.092 | Elargissement: 0.10 nm | T_vib = 2917 |
T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 2 | Score: 0.092 | Elargissement: 0.10 nm | T_vib = 2984 |
T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 3 | Score: 0.092 | Elargissement: 0.10 nm | T_vib = 2984 |
T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 1 | Score: 0.102 | Elargissement: 0.10 nm | T_vib = 4703 |
```

```

T_rot:    712 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.102 | Elargissement:    0.10 nm | T_vib =    4085 |
T_rot:    714 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.102 | Elargissement:    0.10 nm | T_vib =    4080 |
T_rot:    714 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  1 | Score:    0.394 | Elargissement:    0.10 nm | T_vib =   10000 |
T_rot:   1207 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.392 | Elargissement:    0.10 nm | T_vib =    8321 |
T_rot:   1213 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.392 | Elargissement:    0.10 nm | T_vib =    8285 |
T_rot:   1213 K | Scale 1.00000000 | Décalage: 0.0000 nm

```

```

[85]: spectra_simulation_whole = [xs_sim.
    ↪get_whole_spectrum(params_fit_center_pulses[i][5]*W+params_fit_center_pulses[i][4],
    ↪T_el=1_000, T_vib=params_fit_center_pulses[i][1],
    ↪T_rot=params_fit_center_pulses[i][2],
    ↪sigma_exp=params_fit_center_pulses[i][3]) for i in
    ↪range(len(params_fit_center_pulses))]

```

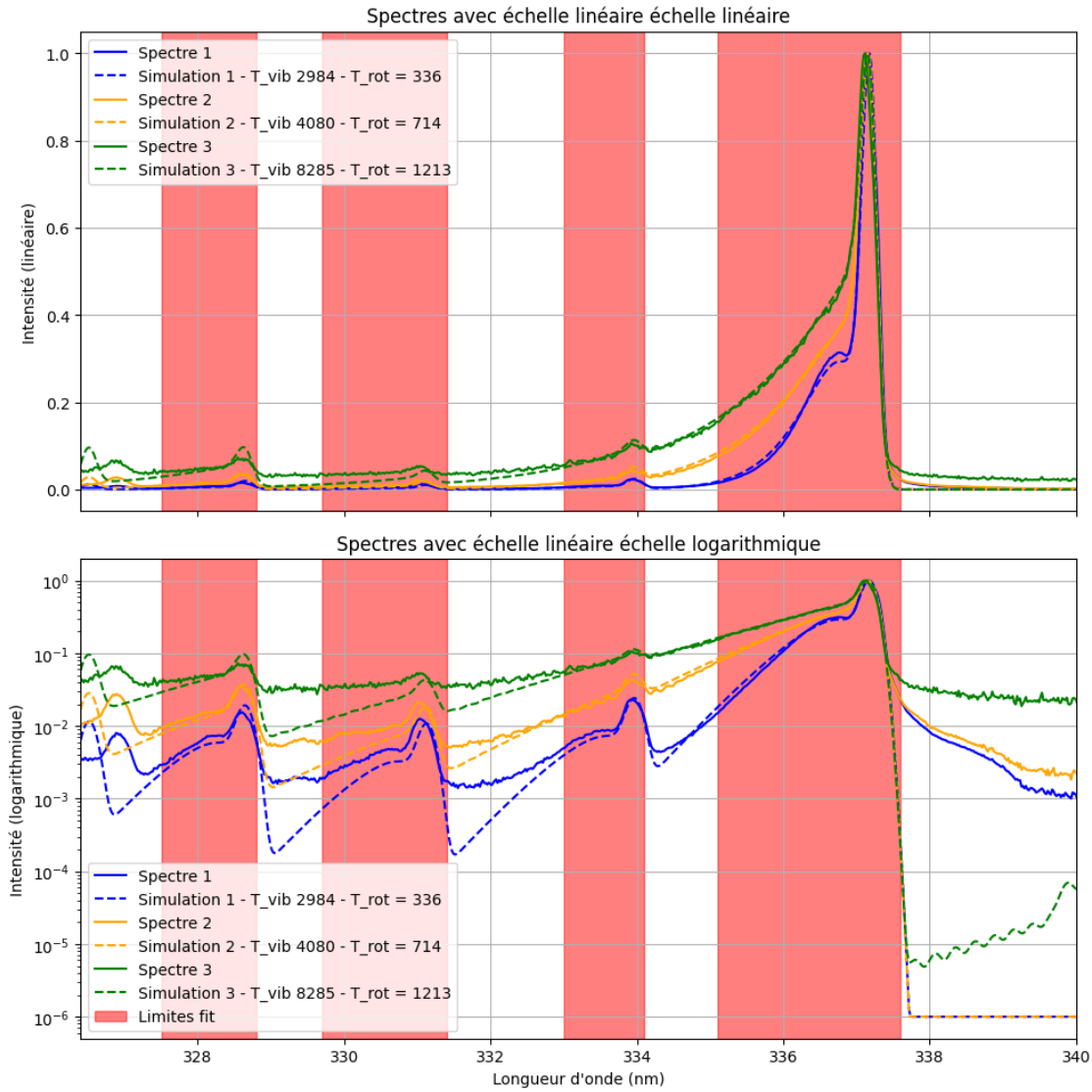
```

[86]: fig, axs = plt.subplots(2, 1, figsize=(10, 10), sharex=True)

# Call the function
plot_spectra_and_simulation(spectra_simulation_whole, axs[0], axs[1],
    ↪fit_areas=areas, simulation_labels = [f'Simulation {i+1} - T_vib {tv:4.0f} -
    ↪T_rot = {tr:3.0f}' for i, (tv, tr) in
    ↪enumerate(zip(params_fit_center_pulses[:,1], params_fit_center_pulses[:
    ↪,2]))], title="Spectres avec échelle linéaire", xlims=(np.min(W), 340),
    ↪epsilon=1e-6)

plt.tight_layout()
plt.show()

```



```
[87]: # Calculer les scores pour chaque valeur de T_rot
T_rot_range = np.linspace(200, 500, 50)
sigma_exp_range = np.linspace(0.05, 0.15, 20)
w_decalage_range = np.linspace(-0.2, 0.3, 20)
T_vib_range = np.linspace(300, 5_000, 20)
w_scale_range = np.linspace(0.999, 1.001, 20)

scores_T_rot = np.zeros_like(T_rot_range)
scores_sigma_exp = np.zeros_like(sigma_exp_range)
scores_w_decalage = np.zeros_like(w_decalage_range)
scores_T_vib = np.zeros_like(T_vib_range)
scores_w_scale = np.zeros_like(w_scale_range)
```

```

for idx, T_rot in enumerate(T_rot_range):
    scores_T_rot[idx] = xs_utils.compute_score_fit(
        c_spectrum_pulses[0],
        xs_sim.
        ↪get_whole_spectrum(params_fit_center_pulses[0][5]*W+params_fit_center_pulses[0][4],
        ↪T_el=1_000, T_vib=params_fit_center_pulses[0][1], T_rot=T_rot,
        ↪sigma_exp=params_fit_center_pulses[0][3])
    )

for idx, sigma_exp in enumerate(sigma_exp_range):
    scores_sigma_exp[idx] = xs_utils.compute_score_fit(
        c_spectrum_pulses[0],
        xs_sim.
        ↪get_whole_spectrum(params_fit_center_pulses[0][5]*W+params_fit_center_pulses[0][4],
        ↪T_el=1_000, T_vib=params_fit_center_pulses[0][1],
        ↪T_rot=params_fit_center_pulses[0][2], sigma_exp=sigma_exp)
    )

for idx, w_decalage in enumerate(w_decalage_range):
    scores_w_decalage[idx] = xs_utils.compute_score_fit(
        c_spectrum_pulses[0],
        xs_sim.get_whole_spectrum(params_fit_center_pulses[0][5]*W+w_decalage,
        ↪T_el=1_000, T_vib=params_fit_center_pulses[0][1],
        ↪T_rot=params_fit_center_pulses[0][2],
        ↪sigma_exp=params_fit_center_pulses[0][3])
    )

for idx, T_vib in enumerate(T_vib_range):
    scores_T_vib[idx] = xs_utils.compute_score_fit(
        c_spectrum_pulses[0],
        xs_sim.
        ↪get_whole_spectrum(params_fit_center_pulses[0][5]*W+params_fit_center_pulses[0][4],
        ↪T_el=1_000, T_vib=T_vib, T_rot=params_fit_center_pulses[0][2],
        ↪sigma_exp=params_fit_center_pulses[0][3])
    )

for idx, w_scale in enumerate(w_scale_range):
    scores_w_scale[idx] = xs_utils.compute_score_fit(
        c_spectrum_pulses[0],
        xs_sim.get_whole_spectrum(w_scale*W+params_fit_center_pulses[0][4],
        ↪T_el=1_000, T_vib=params_fit_center_pulses[0][1],
        ↪T_rot=params_fit_center_pulses[0][2],
        ↪sigma_exp=params_fit_center_pulses[0][3])
    )

# Tracer le graphe
fig, axs = plt.subplots(1, 5, figsize=(18, 6))

```

```

axs[0].plot(T_rot_range, scores_T_rot, marker='o', label="T_rot")
axs[0].set_xlabel("T_rot (K)")
axs[0].set_ylabel("Score")
axs[0].set_title("Score as a function of T_rot")
axs[0].grid()
axs[0].legend()

axs[1].plot(sigma_exp_range, scores_sigma_exp, marker='o', label="sigma_exp",
            color='red')
axs[1].set_xlabel("Élargissement (sigma_exp) (nm)")
axs[1].set_ylabel("Score")
axs[1].set_title("Score as a function of sigma_exp")
axs[1].grid()
axs[1].legend()

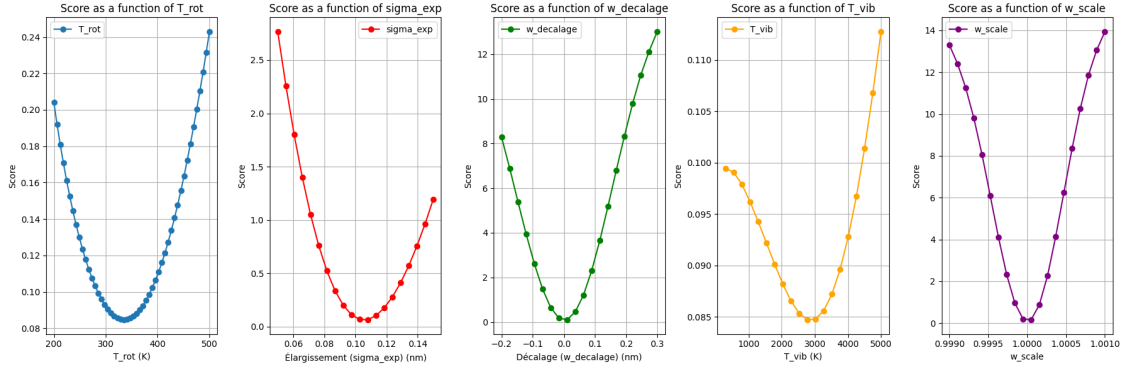
axs[2].plot(w_decalage_range, scores_w_decalage, marker='o',
            label="w_decalage", color='green')
axs[2].set_xlabel("Décalage (w_decalage) (nm)")
axs[2].set_ylabel("Score")
axs[2].set_title("Score as a function of w_decalage")
axs[2].grid()
axs[2].legend()

axs[3].plot(T_vib_range, scores_T_vib, marker='o', label="T_vib",
            color='orange')
axs[3].set_xlabel("T_vib (K)")
axs[3].set_ylabel("Score")
axs[3].set_title("Score as a function of T_vib")
axs[3].grid()
axs[3].legend()

axs[4].plot(w_scale_range, scores_w_scale, marker='o', label="w_scale",
            color='purple')
axs[4].set_xlabel("w_scale")
axs[4].set_ylabel("Score")
axs[4].set_title("Score as a function of w_scale")
axs[4].grid()
axs[4].legend()

plt.tight_layout()
plt.savefig("./res/scores_function.png", dpi=300, bbox_inches='tight')
plt.show()

```



```
[88]: # Récupérons les données pour plus de transitions
# On fait juste pour le centre sinon ça prendrait trop de temps
params_whole_sim = {
    # name : np.array([process_whole_fit(xs_utils.compute_spectra(d)) if
    ↪ is_inside_one_of_the_pulses(delay) else [0]*6 for delay, d in zip(delays,
    ↪ data)]) for name, (delays, data) in plasmas_datas.items())
    name : np.array([process_whole_fit(xs_utils.compute_spectra(d)) if
    ↪ is_inside_one_of_the_pulses(delay) else [0]*6 for delay, d in zip(delays,
    ↪ data)]) if name=="center" else np.zeros((len(delays), 6)) for name, (delays,
    ↪ data) in plasmas_datas.items())
}
```

```
Iteration 1 | Score: 0.121 | Elargissement: 0.10 nm | T_vib = 4015 |
T_rot: 342 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 2 | Score: 0.121 | Elargissement: 0.10 nm | T_vib = 4068 |
T_rot: 341 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 3 | Score: 0.121 | Elargissement: 0.10 nm | T_vib = 4068 |
T_rot: 341 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 1 | Score: 0.092 | Elargissement: 0.10 nm | T_vib = 2917 |
T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 2 | Score: 0.092 | Elargissement: 0.10 nm | T_vib = 2984 |
T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 3 | Score: 0.092 | Elargissement: 0.10 nm | T_vib = 2984 |
T_rot: 336 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 1 | Score: 0.099 | Elargissement: 0.10 nm | T_vib = 3311 |
T_rot: 379 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 2 | Score: 0.099 | Elargissement: 0.10 nm | T_vib = 3336 |
T_rot: 379 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 3 | Score: 0.099 | Elargissement: 0.10 nm | T_vib = 3336 |
T_rot: 379 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 1 | Score: 0.099 | Elargissement: 0.10 nm | T_vib = 3554 |
T_rot: 430 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration 2 | Score: 0.099 | Elargissement: 0.10 nm | T_vib = 3515 |
T_rot: 430 K | Scale 1.00000000 | Décalage: 0.0000 nm
```

```

Iteration  3 | Score:    0.099 | Elargissement:  0.10 nm | T_vib =  3515 |
T_rot:    430 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  1 | Score:    0.093 | Elargissement:  0.10 nm | T_vib =  4432 |
T_rot:    674 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.093 | Elargissement:  0.10 nm | T_vib =  3918 |
T_rot:    675 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.093 | Elargissement:  0.10 nm | T_vib =  3915 |
T_rot:    675 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  1 | Score:    0.102 | Elargissement:  0.10 nm | T_vib =  4703 |
T_rot:    712 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.102 | Elargissement:  0.10 nm | T_vib =  4085 |
T_rot:    714 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.102 | Elargissement:  0.10 nm | T_vib =  4080 |
T_rot:    714 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  1 | Score:    0.131 | Elargissement:  0.10 nm | T_vib =  5470 |
T_rot:    754 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.131 | Elargissement:  0.10 nm | T_vib =  4725 |
T_rot:    756 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.131 | Elargissement:  0.10 nm | T_vib =  4719 |
T_rot:    756 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  1 | Score:    0.394 | Elargissement:  0.10 nm | T_vib = 10000 |
T_rot:   1207 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.392 | Elargissement:  0.10 nm | T_vib =  8321 |
T_rot:   1213 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.392 | Elargissement:  0.10 nm | T_vib =  8285 |
T_rot:   1213 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  1 | Score:    0.383 | Elargissement:  0.10 nm | T_vib = 10000 |
T_rot:   1300 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  2 | Score:    0.380 | Elargissement:  0.10 nm | T_vib =  8175 |
T_rot:   1309 K | Scale 1.00000000 | Décalage: 0.0000 nm
Iteration  3 | Score:    0.380 | Elargissement:  0.10 nm | T_vib =  8127 |
T_rot:   1309 K | Scale 1.00000000 | Décalage: 0.0000 nm

```

1.7.5 Résumé

```

[94]: plt.figure(figsize=(15, 7))

name="center"
for i in range(3):
    filter = get_filter(plasmas_datas[name], i)
    label = r"$T_{vib}$ computed by ratio" if i==0 else None
    plt.plot(plasmas_datas[name][0][filter], t_vib_by_ratio[name][filter], ".
    ↵-", color="purple", label=label)

    label = r"$T_{rot}$ computed with branch R" if i==0 else None

```



```

plt.errorbar(plasmas_datas[name][0][filter], t_rot_analytic[name][filter][:
↪,0], yerr=t_rot_analytic[name][filter][:,1], fmt=".-", color="red",
↪label=label)

label = r"$T_{rot}$ by fit on $\nu'=0 \rightarrow \nu'=0$ " if i==0 else None
plt.plot(plasmas_datas[name][0][filter], t_rot_by_fit_simple[name][filter],
↪".-", color="blue", label=label)

label = r"$T_{rot}$ by fit on all transitions" if i==0 else None
plt.plot(plasmas_datas[name][0][filter], params_whole_sim[name][filter][:
↪,2], ".-", color="orange", label=label)

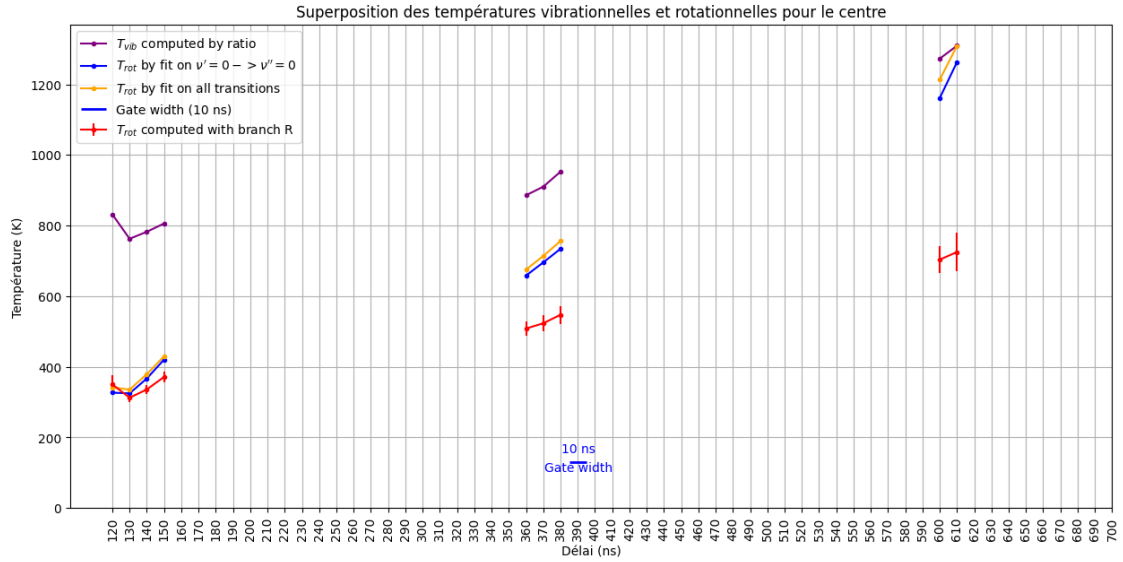
# label = r"$T_{vib}$ by fit on all transitions" if i==0 else None
# plt.plot(plasmas_datas[name][0][filter], params_whole_sim[name][filter][:
↪,1], ".-", color="green", label=label)

# Add a 10 nanoseconds scale bar
M = np.max(t_vib_by_ratio[name])
plt.hlines(y=M * 0.1, xmin=premier_pulse + D_theorique, xmax=premier_pulse +
↪D_theorique + 10, color="blue", linewidth=2, label="Gate width (10 ns)")
plt.text(premier_pulse + D_theorique + 5, M * 0.12, "10 ns", color="blue",
↪ha="center")
plt.text(premier_pulse + D_theorique + 5, M * 0.08, "Gate width", color="blue",
↪ha="center")

plt.ylabel("Température (K)")
plt.xlabel("Délai (ns)")

plt.title("Superposition des températures vibrationnelles et rotationnelles
↪pour le centre")
plt.grid()
plt.xticks(plasmas_datas[name][0], rotation=90)
plt.legend()
plt.ylim(bottom=0)
plt.savefig("./res/superposed_temperatures.png")
plt.show()

```



```
[95]: df = pd.DataFrame({
    "delay": plasmas_datas["center"][0][index_pulses_center],
    "T_vib_by_ratio": t_vib_by_ratio["center"][index_pulses_center],
    "T_rot_by_branch": t_rot_analytic["center"][index_pulses_center,0],
    "T_rot_by_branch-incerti": t_rot_analytic["center"][index_pulses_center,1],
    "T_rot_by_fit_simple": t_rot_by_fit_simple["center"][index_pulses_center],
    "T_rot_by_fit_whole": params_whole_sim["center"][index_pulses_center,2],
    "T_vib_by_fit_whole": params_whole_sim["center"][index_pulses_center,1]
})
df.to_csv("./res/temperatures.csv", index=False)
df
```

```
[95]:
```

	delay	T_vib_by_ratio	T_rot_by_branch	T_rot_by_branch-incerti \
0	130	762.532654	312.498935	11.149773
1	370	910.203186	523.270160	22.108816
2	600	1272.731079	703.482346	38.108269

	T_rot_by_fit_simple	T_rot_by_fit_whole	T_vib_by_fit_whole
0	324.741980	335.538509	2983.605146
1	695.709538	713.772756	4079.850470
2	1161.129945	1213.355147	8284.933923