

## Mission:

To Make the Impossible Possible

### We will:

**Develop** a corporate culture that rewards initiative, enthusiasm and innovation

**Anticipate**, understand and respond to our customers needs quickly

**Lead** in our market space using technology to drive us to the cutting edge of research

**Control** our future through planning, perseverance and commitment to quality



## Products

- Cannabinoids
- Glitazones
- Coelenterazines
- Natural Products
- Steroids
- Nucleosides
- Phosphoramidites
- Fentanyl Salts, Metabolites and Impurity Standards
- Specialty Compounds

**ACCELERATED DRUG  
DISCOVERY AND  
DEVELOPMENT**



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Please visit our website for more information:  
[www.dalton.com](http://www.dalton.com)

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## Why Choose Dalton?

- **One Stop Shop**

Fully integrated chemistry services provider committed to meeting tight timelines

- **Minimize your Burn Rates**

Reduce your cost and shorten your drug development timelines by eliminating the burden of managing multiple contract relationships and tech transfers

- **Flexible Management Team**

Responsive Management with extensive industry experience and flexibility to meet client's needs

- **Convenient Location**

Facility located in the heart of Canada's leading pharma centre



## Services

- **Contract Research**

- **GMP API Manufacturing**

- **GMP Sterile Fill**

Liquid Injectables

Powder Fill

Syringe Fill

- **Custom Synthesis**

Small Molecules

- **Contract Analytical**

Analytical Chemistry

Analytical Development

Microbiology

Stability

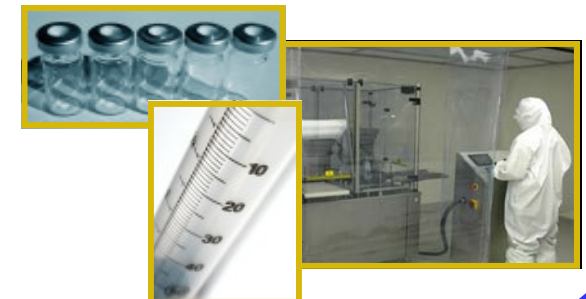


## Company Profile

Dalton is a Health Canada approved contract pharmaceutical manufacturer that supplies chemistry and analytical services to the biotechnology and pharmaceutical industries in the area of chemistry, medicinal chemistry and fine chemical manufacture. Dalton provides cGMP manufacturing and sterile filling services to its customers at any stage of the regulatory process (Phase I, II, III or commercial).

In its 42,000 sq. ft., state of the art cGMP facility, Dalton produces active pharmaceutical ingredients (API) at the gram or kilogram scale. Aseptically filled or terminally sterilized batches of finished drug product in vials or syringes are manufactured under fully validated conditions.

Dalton's analytical laboratory offers method development, validation and ICH stability programs to its clients.



**ACCELERATED**

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